JC05 Rec'd PGT/PTO 1.9 MAR 2002

U.S. DEPARTMENT OF COMMERCE PATENT AND TRADEMARK OFFICE ATTORNEY 'S DOCKET NUMBER FORM PTO-1390 (REV. 11-2000) US 19005 TRANSMITTAL LETTER TO THE UNITED STATES U.S. APPLICATION NO (If known, see 37 CFR 1 5 DESIGNATED/ELECTED OFFICE (DO/EO/US) CONCERNING A FILING UNDER 35 U.S.C. 371 PRIORITY DATE CLAIMED INTERNATIONAL APPLICATION NO. INTERNATIONAL FILING DATE September 22, 1999 PCT/EP00/09111 September 15, 2000 TITLE OF INVENTION CATALYST SYSTEM AND PROCESS FOR THE POLYMERIZATION OF OLEFINS APPLICANT(S) FOR DO/EO/US BASELL TECHNOLOGY COMPANY BV Applicant herewith submits to the United States Designated/Elected Office (DO/EO/US) the following items and other information: 1. X This is a FIRST submission of items concerning a filing under 35 U.S.C. 371. 2. This is a SECOND or SUBSEQUENT submission of items concerning a filing under 35 U.S.C. 371. 3. X This is an express request to begin national examination procedures (35 U.S.C. 371(f)). The submission must include items (5), (6), (9) and (21) indicated below. 4. X The US has been elected by the expiration of 19 months from the priority date (Article 31). 5. X A copy of the International Application as filed (35 U.S.C. 371(c)(2)) a. is attached hereto (required only if not communicated by the International Bureau). x has been communicated by the International Bureau. is not required, as the application was filed in the United States Receiving Office (RO/US). 6. An English language translation of the International Application as filed (35 U.S.C. 371(c)(2)). is attached hereto. has been previously submitted under 35 U.S.C. 154(d)(4). b. 7. X Amendments to the claims of the International Aplication under PCT Article 19 (35 U.S.C. 371(c)(3)) are attached hereto (required only if not communicated by the International Bureau). have been communicated by the International Bureau. have not been made; however, the time limit for making such amendments has NOT expired. have not been made and will not be made. 8. An English language translation of the amendments to the claims under PCT Article 19 (35 U.S.C. 371 (c)(3)). 9. X An oath or declaration of the inventor(s) (35 U.S.C. 371(c)(4)). 10. An English lanugage translation of the annexes of the International Preliminary Examination Report under PCT Article 36 (35 U.S.C. 371(c)(5)). Items 11 to 20 below concern document(s) or information included: An Information Disclosure Statement under 37 CFR 1.97 and 1.98. 11. An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31 is included. 12. x A FIRST preliminary amendment. 13. X A SECOND or SUBSEQUENT preliminary amendment. 14. A substitute specification. 15. A change of power of attorney and/or address letter. 16. A computer-readable form of the sequence listing in accordance with PCT Rule 13ter.2 and 35 U.S.C. 1.821 - 1.825. 17. A second copy of the published international application under 35 U.S.C. 154(d)(4). 18. A second copy of the English language translation of the international application under 35 U.S.C. 154(d)(4). 19. Receipt acknowledgment card Other items or information: 20. X Express Mailing Certificate

			JC10 Rec	4.7577710 1°9	MAR 2002
U.S. APP IC PON/10 PARE	BY ARRIVE IN	TERNATIONAL APPLICATION NO	- aillu thail thail th		KETNUNGER IN ILILE 005
		CT/EP00/09111		CALCULATIONS	
21. The following	ng fees are submitted:	(4) (5)	ł		
BASIC NATIONAL	FEE (37 CFR 1.492 (a)	(1) - (5)): on fee (27 CEP 1 482)			
	al preliminary examinati arch fee (37 CFR 1.445)	ai())) baid to USPTU			
and International Se	arch Report not prepare	d by the EPO or JPO	\$1000.00		
International prelim	inary examination fee (37 CFR 1.482) not paid to			•
USPTO but Interna	tional Search Report pro	epared by the EPO or JPO	\$860.00		
International prelim but international sea	iinary examination fee (arch fee (37 CFR 1.445(37 CFR 1.482) not paid to a)(2)) paid to USPTO	USPTO \$710.00		
but all claims did no	ot satisfy provisions of I	37 CFR 1.482) paid to US PCT Article 33(1)-(4)	3050.00		
International prelim	ninary examination fee (37 CFR 1.482) paid to US	PTO \$100.00		
and all claims satis	fied provisions of PCT A	Article 33(1)-(4)	S100.00	s 890.00	i
				3 0,0.00	
Surcharge of \$130.0 months from the ear	0 for furnishing the oath liest claimed priority dat	or declaration later than te (37 CFR 1.492(e)).	20 30	\$	<u> </u>
CLAIMS	NUMBER FILED	NUMBER EXTRA	RATE	\$	<u></u>
Total claims	27 - 20 =	7	x \$18.00	\$ 126	
Independent claims	1 -3 =	0	x \$80.00	\$ 0	
	DENT CLAIM(S) (if ap	plicable)	+ \$270.00	\$ 0	
	TOTAL	OF ABOVE CALCU	LATIONS =	\$ 1016	
Applicant claim	is small entity status. So	ee 37 CFR 1.27. The fees	indicated above	\$	
are reduced by	1/2.		+	s 1016	
			UBTOTAL =	\$ 1016	
Processing fee of \$1 months from the ear	30.00 for furnishing the liest claimed priority da			\$	
		TOTAL NATIO		\$ 1016	
Fee for recording the accompanied by an	e enclosed assignment (appropriate cover sheet	37 CFR 1.21(h)). The ass (37 CFR 3.28, 3.31). \$40	ignment must be .00 per property +	\$ 40	
		TOTAL FEES E	NCLOSED =	\$ 1056	
				Amount to be refunded:	\$
				charged:	\$
a. A check in	the amount of \$	to cover t	he above fees is enclo	osed.	
			n the amount of $\frac{10}{2}$		he above fees.
A duplica	arge my Deposit Account te copy of this sheet is e	nclosed.			
c. X The Comi	missioner is hereby auth ent to Deposit Account	orized to charge any addit No. <u>08-2336</u> . A dupl	ional fees which may icate copy of this shee	be required, or credit et is enclosed.	any
d. Fees are to informati	o be charged to a credit on should not be include	card. WARNING: Informated on this form. Provide	mation on this form me credit card informati	ay become public. Con and authorization	redit card on PTO-2038.
NOTE: Where a	n appropriate time lim	it under 37 CFR 1.494 o ed to restore the applica	r 1.495 has not been tion to pending statu	met, a petition to res	vive (37 CFR
SEND ALL CORRES			-	ueW. Sa	tterson.
	Patterson		SIGNAT	TURE.	
1	orth America In	с.	Joa	nne W. Patter	son
	tual Property		NAME		
	eton Road			217	
Elkton, N				217	
	10-996- 1658		REGIST	TRATION NUMBER	

fax: 410-996-1560

JC10 Rec'd PCT/PTO 1 9 MAR 2002

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re patent application of)
Jan F. Van Baar et al.)
Serial No. Unknown)
Filed March 19, 2002))
PCT International Appln. PCT/EP00/09111))
Filed September 15, 2000)
For CATALYST SYSTEM AND PROCESS FOR THE POLYMERIZATION OF OLEFINS))

Commissioner for Patents Box PCT Washington, D. C. 20231

Sir:

PRELIMINARY AMENDMENT

This is a preliminary amendment to US 19005, filed herewith. US 19005 is the U.S. national phase of International Application PCT/EP00/09111, filed September 15, 2000.

International Application PCT/EP00/09111 was published under PCT Article 21(a) in English.

Please enter the amendments listed below in the above-identified application.

In the Specification

In the first sentence after the title, please insert:

--This application is the U.S. national phase of International Application PCT/EP00/09111, filed September 15, 2000.--

In the Claims

- 1. (Amended) A catalyst system for the polymerization of olefins comprising the product obtained by contacting the following components:
- (A) a metallocene complex of formula (I):

$$(Cp)(ZR_m^1)_n(A)_rML_p$$
 (I)

wherein $(ZR_m^1)_n$ is a divalent group bridging Cp and A, Z being C, Si, Ge, N or P, and the R^1 groups, equal or different from each other, being selected from the group consisting of (1) hydrogen and (2) linear or branched, saturated or unsaturated C_1 - C_{20} alkyl, C_3 - C_{20} cycloalkyl, C_6 - C_{20} aryl, C_7 - C_{20} alkylaryl or C_7 - C_{20} arylalkyl groups;

Cp is a substituted or unsubstituted cyclopentadienyl group, optionally condensed to one or more substituted or unsubstituted, saturated, unsaturated or aromatic rings, containing from 4 to 6 carbon atoms, optionally containing one or more heteroatoms;

A is -O-, -S-, -N(R^2)-, wherein R^2 is hydrogen, a linear or branched, saturated or unsaturated C_1 - C_{20} alkyl, C_3 - C_{20} cycloalkyl, C_6 - C_{20} aryl, C_7 - C_{20} alkylaryl or C_7 - C_{20} arylalkyl, or A has the same meaning as Cp;

M is a transition metal belonging to group 3, 4, 5, 6, the lanthanide, or the actinide groups of the Periodic Table;

the substituents L, same or different from each other, are monoanionic sigma ligands selected from the group consisting of hydrogen, halogen, $-R^3$, $-OR^3$, $-OCOR^3$, $-SR^3$, $-NR^3$ ₂ and $-PR^3$ ₂, wherein R^3 is a linear or branched, saturated or unsaturated C_1 - C_{20} alkyl, C_3 - C_{20} cycloalkyl, C_6 - C_{20} aryl, C_7 - C_{20} alkylaryl or C_7 - C_{20} arylalkyl group, optionally containing one or more Si or Ge atoms;

m is 1 or 2; n is an integer ranging from 0 to 4; r is 0 or 1; n is 0 when r is 0; p is an integer equal to the oxidation state of the metal M minus 2 when r=1, and minus 1 when r=0, and ranges from 1 to 4;

(B) an organometallic aluminium compound of formula (II):

Al
$$[CH_2-C(Ar)R^4R^5]_xH_y$$
 (II)

wherein Ar is a substituted or unsubstituted aryl group having from 6 to 20 carbon atoms;

 R^4 is a linear or branched, saturated or unsaturated, C_1 - C_{10} alkyl, C_6 - C_{20} aryl, C_7 - C_{20} arylalkyl or C_7 - C_{20} alkylaryl; R^5 is selected from the group consisting of (1) hydrogen and (2) a linear or branched, saturated or unsaturated, C_1 - C_{10} alkyl, C_6 - C_{20} aryl, C_7 - C_{20} arylalkyl or C_7 - C_{20} alkylaryl group; the substituents R^4 and R^5 optionally form a ring, having 3 to 8 carbon atoms; a carbon atom in the compound of formula (II) being optionally replaced by a Si or a Ge atom;

 \mathbf{x} is 2 or 3; $\mathbf{y} = 3 - \mathbf{x}$; and

(C) water;

the molar ratio between said organometallic aluminium compound (B) and said water (C) being between 1:1 and 100:1.

- 3. (Amended) The catalyst system according to claim 1 wherein the molar ratio between said organometallic aluminium compound (B) and said metallocene complex (A) ranges from 50 to 50,000.
- 9. (Amended) The catalyst system according to claim 1 wherein, in said organometallic aluminium compound of formula (II), Ar corresponds to formula (III):

$$R^{6}$$
 R^{7}
 R^{8}
(III)

wherein R^6 , R^8 and R^{10} are selected from the group consisting of hydrogen, halogen, $-R^3$, $-C(O)R^3$, $-OR^3$, $-SR^3$, $-NR^3$ ₂ and $-NO_2$, wherein R^3 has the meaning reported in claim 1; the groups R^7 and R^9 are selected from the group consisting of (1) hydrogen, (2) halogen, and (3) linear or branched, saturated or unsaturated C_1 - C_{20} alkyl, C_3 - C_{20} cycloalkyl, C_6 - C_{20} aryl, C_7 - C_{20} alkylaryl and C_7 - C_{20} arylalkyl groups, optionally containing one or more Si or Ge atoms; and two adjacent substituents R^6 - R^{10} optionally form a ring having 3 to 8 carbon atoms;

R⁴ is a C₁-C₅ alkyl group; and

R⁵ is hydrogen or a C₁-C₅ alkyl group.

- 12. (Amended) A catalyst system for the polymerization of olefins comprising the product obtained by contacting the following components:
 - (A) a metallocene complex of formula (I):

$$(Cp)(ZR_m^1)_n(A)_rML_p$$
 (I)

wherein M, Cp, (ZR¹_m)_n, A, L, r and p have the meanings reported in claim 1; and

(B') the product of the reaction between water and an organometallic aluminium compound of formula (II):

Al
$$[CH_2-C(Ar)R^4R^5]_xH_y$$
 (II)

wherein Ar, R⁴, R⁵, x and y have the meanings reported in claim 1;

the molar ratio between said organometallic aluminium compound and said water being between 1:1 and 100:1.

13. (Amended) The catalyst system according to claim 1, wherein said metallocene complex is pre-alkylated with one or more organometallic aluminium compounds of formula (IV):

$$AlR^{11}_{3-z}H_z$$
 (IV)

wherein R^{11} is a linear or branched, saturated or unsaturated, C_1 - C_{10} alkyl, C_6 - C_{20} aryl, C_7 - C_{20} arylalkyl or C_7 - C_{20} alkylaryl and ${\bf z}$ is 0 or 1.

14. (Amended) An alumoxane obtained by contacting an organometallic aluminium compound of formula (II)

Al
$$[CH_2-C(Ar)R^4R^5]_xH_y$$
 (II)

wherein Ar, R⁴, R⁵, x and y have the meanings reported in claim 1, with water, the molar ratio between said organometallic aluminium compound and said water being between 1:1 and 100:1.

15. (Amended) A process for the polymerization of one or more olefins in the presence of a catalyst system according to claim 1.

In the Abstract

Delete the last paragraph and insert in its place:

--the molar ratio (B)/(C) being between 1:1 and 100:1. These catalysts show an improved activity with respect to known catalysts, wherein different aluminium compounds are used.--

Add the following new claims:

19. (New) The catalyst system according to claim 12, wherein said metallocene complex is pre-alkylated with one or more organometallic aluminium compounds of formula (IV):

$$AlR^{11}_{3-z}H_z$$
 (IV)

wherein R^{11} is a linear or branched, saturated or unsaturated, C_1 - C_{10} alkyl, C_6 - C_{20} aryl, C_7 - C_{20} arylalkyl or C_7 - C_{20} alkylaryl and ${\bf z}$ is 0 or 1.

- 20. (New) A process for the polymerization of one or more olefins in the presence of a catalyst system according to claim 12.
- 21. (New) A process for the polymerization of one or more olefins in the presence of a catalyst system according to claim 13.
- 22. (New) The process according to claim 12, wherein said olefin is an α -olefin having the formula CH₂=CHR, wherein R is hydrogen or a C₁-C₂₀ alkyl radical.
- 23. (New) The process according to claim 13 wherein said olefin is an α -olefin having the formula CH₂=CHR, wherein R is hydrogen or a C₁-C₂₀ alkyl radical.
- 24. (New) The process according to claim 12, wherein said α-olefin is selected from the group consisting of propylene, 1-butene, 4-methyl-1-pentene, 1-hexene and 1-octene.
- 25. (New) The process according to claim 13, wherein said α -olefin is selected from the group consisting of propylene, 1-butene, 4-methyl-1-pentene, 1-bexene and 1-octene.
- 26. (New) The process according to claim 12, wherein ethylene is copolymerized with an α -olefin having the formula CH₂=CHR', wherein R' is a linear, branched or cyclic C₁-C₂₀ alkyl radical, or with a cycloolefin, and optionally with a polyene.
- 27. (New) The process according to claim 13, wherein ethylene is copolymerized with an α -olefin having the formula CH₂=CHR', wherein R' is a linear, branched or cyclic C₁-C₂₀ alkyl radical, or with a cycloolefin, and optionally with a polyene.

REMARKS

A sentence indicating that the present invention is the U.S. national phase of a PCT application has been inserted in the specification.

Claims 1, 3, 9, and 12-15 have been amended to eliminate multiple dependent claims and to put the claims in the proper format for filing.

The abstract has been amended to remove a superfluous word.

New claims 19-27 have been added.

Separate pages entitled "Version with Markings to Show Changes Made" are attached to indicate the amendments made to the above-identified application.

Respectfully submitted,

Joanne W. Patterson Registration No. 31,217 Agent for Applicants

Basell North America Inc. 912 Appleton Road Elkton, MD 21921

March 19, 2002

Agent's Telephone No.: 410-996-1658

I hereby certify that this correspondence is being deposited with the United States Postal Service as "Express Mail Post Office to Addressee" in an envelope addressed to: Box PCT, Commissioner for Patents, Washington, DC 20231 on March 19, 2002 with the number of the Express Mail label being

EK794125403US

VERSION WITH MARKINGS TO SHOW CHANGES MADE

In the Claims

- 1. (Amended) A catalyst system for the polymerization of olefins comprising the product obtained by contacting the following components:
- (A) a metallocene complex of formula (I):

$$(Cp)(ZR^{1}_{m})_{n}(A)_{r}ML_{p}$$
 (I)

wherein (ZR¹_m)_n is a divalent group bridging Cp and A, Z being C, Si, Ge, N or P, and the R¹ groups, equal or different from each other, being <u>selected from the group consisting of (1)</u> hydrogen [or] <u>and (2)</u> linear or branched, saturated or unsaturated C₁-C₂₀ alkyl, C₃-C₂₀ cycloalkyl, C₆-C₂₀ aryl, C₇-C₂₀ alkylaryl or C₇-C₂₀ arylalkyl groups;

Cp is a substituted or unsubstituted cyclopentadienyl group, optionally condensed to one or more substituted or unsubstituted, saturated, unsaturated or aromatic rings, containing from 4 to 6 carbon atoms, optionally containing one or more heteroatoms;

A is -O-, -S-, -N(\mathbb{R}^2)-, wherein \mathbb{R}^2 is hydrogen, a linear or branched, saturated or unsaturated C_1 - C_{20} alkyl, C_3 - C_{20} cycloalkyl, C_6 - C_{20} aryl, C_7 - C_{20} alkylaryl or C_7 - C_{20} arylalkyl, or A has the same meaning [of] <u>as</u> Cp;

M is a transition metal belonging to group 3, 4, 5, 6, [or to] the lanthanide, or the actinide groups of the Periodic Table;

the substituents L, same or different from each other, are monoanionic sigma ligands selected from the group consisting of hydrogen, halogen, $-R^3$, $-OR^3$, $-OCOR^3$, $-SR^3$, $-NR^3_2$ and $-PR^3_2$, wherein R^3 is a linear or branched, saturated or unsaturated C_1 - C_{20} alkyl, C_3 - C_{20} cycloalkyl, C_6 - C_{20} aryl, C_7 - C_{20} alkylaryl or C_7 - C_{20} arylalkyl group, optionally containing one or more Si or Ge atoms;

m is 1 or 2; n is an integer ranging from 0 to 4; r is 0 or 1; n is 0 when r is 0; p is an integer equal to the oxidation state of the metal M minus 2 when r=1, and minus 1 when r=0, and ranges from 1 to 4;

(B) an organometallic aluminium compound of formula (II):

Al
$$[CH_2-C(Ar)R^4R^5]_xH_y$$
 (II)

wherein Ar is a substituted or unsubstituted aryl group having from 6 to 20 carbon atoms:

 R^4 is a linear or branched, saturated or unsaturated, C_1 - C_{10} alkyl, C_6 - C_{20} aryl, C_7 - C_{20} arylalkyl or C_7 - C_{20} alkylaryl; R^5 is selected from the group consisting of (1) hydrogen [or] and (2) a linear or branched, saturated or unsaturated, C_1 - C_{10} alkyl, C_6 - C_{20} aryl, C_7 - C_{20} arylalkyl or C_7 - C_{20} alkylaryl group; the substituents R^4 and R^5 optionally form a ring, having 3 to 8 carbon atoms; a carbon atom in the compound of formula (II) being optionally replaced by a Si or a Ge atom;

x is 2 or 3; y = 3-x; and

(C) water;

the molar ratio between said organometallic aluminium compound (B) and said water (C) being [comprised] between 1:1 and 100:1.

- 3. (Amended) The catalyst system according to claim 1 [or 2]. Wherein the molar ratio between said organometallic aluminium compound (B) and said metallocene complex (A) ranges from 50 to 50,000.
- 9. (Amended) The catalyst system according to claim 1 wherein, in said organometallic aluminium compound of formula (II), Ar corresponds to formula (III):

$$R^{6}$$
 R^{7}
 R^{8}
(III)

wherein R⁶, R⁸ and R¹⁰ are selected from the group consisting of hydrogen, halogen, -R³, -C(O)R³, -OR³, -SR³, -NR³₂ and -NO₂, wherein R³ has the meaning reported in claim 1; the groups R⁷ and R⁹ are selected from the group consisting of (1) hydrogen, (2) halogen, and (3) linear or branched, saturated or unsaturated C₁-C₂₀ alkyl, C₃-C₂₀ cycloalkyl, C₆-C₂₀ aryl, C₇-C₂₀ alkylaryl and C₇-C₂₀ arylalkyl groups, optionally containing one or more Si or Ge atoms; and two adjacent substituents R⁶-R¹⁰ optionally form a ring[,] having 3 to 8 carbon atoms;

R⁴ is a C₁-C₅ alkyl group; and

 R^5 is hydrogen or a C_1 - C_5 alkyl group.

- 12. (Amended) A catalyst system for the polymerization of olefins comprising the product obtained by contacting the following components:
 - (A) a metallocene complex of formula (I):

$$(Cp)(ZR^{1}_{m})_{n}(A)_{r}ML_{p} \qquad (I)$$

wherein M, Cp, (ZR¹_m)_n, A, L, r and p have the meanings reported in claim 1; and

(B') the product of the reaction between water and an organometallic aluminium compound of formula (II):

Al
$$[CH_2-C(Ar)R^4R^5]_xH_y$$
 (II)

wherein Ar, R⁴, R⁵, x and y have the meanings reported in claim 1;

the molar ratio between said organometallic aluminium compound and said water being [comprised] between 1:1 and 100:1.

13. (Amended) The catalyst system according to claim 1 [or 12], wherein said metallocene complex is pre-alkylated with one or more organometallic aluminium compounds of formula (IV):

$$AlR^{11}_{3-z}H_z$$
 (IV)

wherein R^{11} is a linear or branched, saturated or unsaturated, C_1 - C_{10} alkyl, C_6 - C_{20} aryl, C_7 - C_{20} arylalkyl or C_7 - C_{20} alkylaryl and \mathbf{z} is 0 or 1.

14. (Amended) An alumoxane obtained by contacting an organometallic aluminium compound of formula (II)

Al
$$[CH_2-C(Ar)R^4R^5]_xH_y$$
 (II)

wherein Ar, R⁴, R⁵, x and y have the meanings reported in claim 1, with water, the molar ratio between said organometallic aluminium compound and said water being [comprised] between 1:1 and 100:1.

15. (Amended) A process for the polymerization of one or more olefins in the presence of a catalyst system according to <u>claim 1</u>[any of claims 1 to 13].

In the Abstract

Delete the last paragraph and insert in its place:

--the molar ratio (B)/(C) being [comprised] between 1:1 and 100:1. These catalysts show an improved activity with respect to known catalysts, wherein different aluminium compounds are used.--



CATALYST SYSTEM AND PROCESS FOR THE POLYMERIZATION OF OLEFINS FIELD OF THE INVENTION

The present invention relates to a high activity catalyst system for the polymerization of olefins comprising the reaction product of a metallocene complex, a specific organometallic aluminium compound and water.

The invention also relates to a processes for the polymerization of olefins carried out in the presence of said catalyst system.

PRIOR ART DISCLOSURE

Homogeneous catalytic systems based on metallocene complexes activated by alumoxanes are known to be active in the polymerization of olefins since mid-1980s; the class of alkylalumoxanes, commonly obtained by reacting trialkylaluminium compound and water in a molar ratio of 1:1 to 100:1, comprises oligomeric linear and/or cyclic alumoxanes represented by the formulae:

$$R$$
 Al O $(Al$ $O)_n$ Al R

for linear oligomeric alumoxanes, and

for cyclic oligomeric alumoxanes, wherein the substituents R can be alkyl, alkenyl or alkylaryl radicals, having 1-20 and preferably 1-8 carbon atoms, **n** ranges from 1 to 40, preferably from 10 to 20, and **m** ranges from 3 to 40, preferably from 3 to 20.

In the most widely used alumoxanes, the R substituents are methyl, ethyl or isobutyl groups; methylalumoxane is the most common cocatalyst in metallocene catalyst systems.

Nevertheless alkylalumoxanes, and in particular methylalumoxane, though very active in metallocene-based catalyst systems, exhibit several inherent problems in use, such as the need for high alumoxane/metallocene molar ratios to produce satisfactory catalytic activities, their high reactivity toward impurities (moisture, alcohols etc.) and their easy flammability; accordingly, some of the developments in this area involved a search for alternative cocatalysts. EP-A-0,575,875, in the name of the same Applicant, describes homogeneous catalytic systems for the polymerization of olefins comprising the product obtained by contacting a

metallocene complex of Ti, Zr or Hf, an organometallic aluminium compound wherein at least one alkyl is different from a straight alkyl, and water; in the working examples, said organometallic aluminium compound is triisobutylaluminium or triisohexylaluminium.

WO 96/02580, in the name of the same Applicant, describes improved catalytic systems comprising a metallocene compound and the product obtained by contacting water and an organometallic aluminium compound of formula:

$$Al(CH_2-CR^4R^5R^6)_wR^7_vH_z$$

wherein R^4 is a C_1 - C_{10} alkyl, alkenyl or arylalkyl group; R^5 is a C_3 - C_{50} hydrocarbon different from a straight alkyl or alkenyl group; R^6 is hydrogen or a C_1 - C_{10} hydrocarbon group; R^7 is a C_1 - C_{10} hydrocarbon group; w is 1-3; z is 0 or 1; and y=3-w-z.

WO 99/21899, in the name of the same Applicant, describes a catalyst system for olefin polymerization comprising a metallocene compound activated by the product obtained by contacting water and an organometallic aluminium compound of formula:

$$Al(CH_2-CR^4R^5-CR^6R^7R^8)_wR_a^9H_z$$

wherein R⁴ is a C₁-C₁₀ alkyl or C₇-C₂₀ arylalkyl group; R⁵ is hydrogen, a C₁-C₁₀ alkyl or a C₇-C₁₀ arylalkyl group; R⁶ and R⁷ are C₁-C₁₀ hydrocarbon groups; R⁸ is hydrogen or a C₁-C₁₀ hydrocarbon group; R⁹ is a C₁-C₁₀ alkyl or C₇-C₁₀ arylalkyl group; w is 1-3; z is 0 or 1; and q=3-w-z. Said organometallic aluminium compounds are characterized by presenting one or more (CH₂-CR⁴R⁵-CR⁶R⁷R⁸) groups, which are alkyl groups substituted in beta and gamma positions; illustrative examples of these compounds are tris(2,3-dimethyl-hexyl)aluminium, tris(2,3,3,-trimethyl-butyl)aluminium, tris(2-methyl-3-ethyl-pentyl)aluminium and tris(2-ethyl-3-methyl-pentyl)aluminium.

However, the catalysts described in the above-cited patent applications exert activities in the polymerization of olefins which are not completely satisfactory in all the applications. Therefore, the problem addressed by the present invention is to further improve the activities of the above-discussed known catalysts.

To solve this problem, novel catalysts have unexpectedly been found which are suitable for the polymerization of olefins and possess a considerably improved activity compared with the known catalysts.

SUMMARY OF THE INVENTION

The Applicant has now found a catalyst system for the polymerization of olefins comprising the

PCT/EP00/09111

WO 01/21674

product obtained by contacting the following components:

(A) a metallocene complex of formula (I):

$$(Cp)(ZR_{m}^{1})_{n}(A)_{r}ML_{p} \qquad (I)$$

wherein $(ZR_{m}^{1})_{n}$ is a divalent group bridging Cp and A, Z being C, Si, Ge, N or P, and the R^{1} groups, equal or different from each other, being hydrogen or linear or branched, saturated or unsaturated C_{1} - C_{20} alkyl, C_{3} - C_{20} cycloalkyl, C_{6} - C_{20} aryl, C_{7} - C_{20} alkylaryl or C_{7} - C_{20} arylalkyl groups;

Cp is a substituted or unsubstituted cyclopentadienyl group, optionally condensed to one or more substituted or unsubstituted, saturated, unsaturated or aromatic rings, containing from 4 to 6 carbon atoms, optionally containing one or more heteroatoms;

A is -O-, -S-, -N(R^2)-, wherein R^2 is hydrogen, a linear or branched, saturated or unsaturated C_1 - C_{20} alkyl, C_3 - C_{20} cycloalkyl, C_6 - C_{20} aryl, C_7 - C_{20} alkylaryl or C_7 - C_{20} arylalkyl, or A has the same meaning of Cp;

M is a transition metal belonging to group 3, 4, 5, 6 or to the lanthanide or actinide groups of the Periodic Table of the Elements (IUPAC version);

the substituents L, same or different from each other, are monoanionic sigma ligands selected from the group consisting of hydrogen, halogen, $-R^3$, $-OR^3$, $-OCOR^3$, $-SR^3$, $-NR^3$ ₂ and $-PR^3$ ₂, wherein R^3 is a linear or branched, saturated or unsaturated C_1 - C_{20} alkyl, C_3 - C_{20} cycloalkyl, C_6 - C_{20} aryl, C_7 - C_{20} alkylaryl or C_7 - C_{20} arylalkyl group, optionally containing one or more Si or Ge atoms; preferably, the substituents L are the same;

m is 1 or 2, and more specifically it is 1 when Z is N or P, and it is 2 when Z is C, Si or Ge; n is an integer ranging from 0 to 4;

 \mathbf{r} is 0 or 1; \mathbf{n} is 0 when \mathbf{r} is 0;

p is an integer equal to the oxidation state of the metal M minus 2 when r=1, and minus 1 when r=0, and ranges from 1 to 4;

(B) an organometallic aluminium compound of formula (II):

Al
$$[CH_2-C(Ar)R^4R^5]_xH_y$$
 (II)

wherein

Ar is a substituted or unsubstituted aryl group having from 6 to 20 carbon atoms;

 R^4 is a linear or branched, saturated or unsaturated, C_1 - C_{10} alkyl, C_6 - C_{20} aryl, C_7 - C_{20} arylalkyl or C_7 - C_{20} alkylaryl; R^5 is hydrogen or a linear or branched, saturated or unsaturated, C_1 - C_{10}

alkyl, C_6 - C_{20} aryl, C_7 - C_{20} arylalkyl or C_7 - C_{20} alkylaryl group; the substituents R^4 and R^5 optionally form a ring, having 3 to 8 carbon atoms;

a carbon atom in the compound of formula (II) being optionally replaced by a Si or a Ge atom;

x is 2 or 3; y = 3-x; and

(C) water;

the molar ratio between the organometallic aluminium compound (B) and water (C) being comprised between 1:1 and 100:1,

DETAILED DESCRIPTION OF THE INVENTION

In the catalyst system according to the invention, the molar ratio between the organometallic aluminium compound (B) and water (C) is comprised between 1:1 to 100:1, and preferably between 1:1 and 50:1. A suitable value for the Al/H₂O molar ratio is about 2.

The molar ratio between the organometallic aluminium compound (B) and the metallocene complex (A), calculated as Al/M molar ratio, preferably ranges from 50 to 50000, and more preferably from 500 to 5000.

In the metallocene complex (A) of formula (I), the divalent bridge $(ZR_m^1)_n$ is preferably selected from the group consisting of CR_2^1 , $(CR_2^1)_2$, $(CR_2^1)_3$, SiR_2^1 , GeR_2^1 , NR_2^1 and PR_2^1 , R_2^1 having the meaning reported above; more preferably, said divalent bridge is $Si(CH_3)_2$, $SiPh_2$, CH_2 , $(CH_2)_2$, $(CH_2)_3$ or $C(CH_3)_2$.

The variable **m** is 1 or 2; the variable **n** ranges from 0 to 4 and, when n > 1, the atoms Z can be the same or different from each other, such as in divalent bridges $-CH_2-O_-$, $-CH_2-S_-$ and $-CH_2-S_ -CH_3-S_-$ and $-CH_3-S_ -CH_3-S_-$ and $-CH_3-S_-$ and

The ligand Cp, which is π-bonded to said metal M, is preferably selected from the group consisting of cyclopentadienyl, mono-, di-, tri- and tetra-methyl cyclopentadienyl; 4-butyl-cyclopentadienyl; 4-adamantyl-cyclopentadienyl; indenyl; mono-, di-, tri- and tetra-methyl indenyl; 3-butyl-indenyl; 3-trimethylsilyl-indenyl; 4,5,6,7-tetrahydroindenyl; fluorenyl; 5,10-dihydroindeno[1,2-b]indol-10-yl; N-methyl- or N-phenyl-5,10-dihydroindeno [1,2-b]indol-10-yl; 5,6-dihydroindeno[2,1-b]indol-6-yl; N-methyl-or N-phenyl-5,6-dihydroindeno[2,1-b]indol-6-yl; azapentalene-4-yl; thiapentalene-4-yl; thiapentalene-6-yl; thiapentalene-6-yl; mono-, di- and tri-methyl-azapentalene-4-yl.

The group A is -O-, -S-, -N(R2)-, wherein R2 is hydrogen, a linear or branched, saturated or

PCT/EP00/09111

WO 01/21674

unsaturated C_1 - C_{20} alkyl, C_3 - C_{20} cycloalkyl, C_6 - C_{20} aryl, C_7 - C_{20} alkylaryl or C_7 - C_{20} arylalkyl, or A has the same meaning of Cp.

When A is $-N(R^2)$ -, a suitable class of metallocene complexes (A) for use in the catalysts complexes of the invention comprises the well-known constrained geometry catalysts, as described in EP-A-0 416 815, EP-A-0 420 436, EP-A-0 671 404, EP-A-0 643 066 and WO 91/04257.

According to a preferred embodiment of the invention, the group A has the same meaning of Cp, and it is preferably cyclopentadienyl, indenyl or tetrahydroindenyl.

Suitable metallocene complexes (A) that may be used in the catalysts system according to the present invention are described in USP 5,324,800 and EP-A-0 129 368.

The metal M is preferably Ti, Zr or Hf, and more preferably Zr.

The substituents L are preferably the same and are selected from the group consisting of halogens, R³, -OR³ and -NR³₂; wherein R³ is a C₁-C₇ alkyl, C₆-C₁₄ aryl or C₇-C₁₄ arylalkyl group, optionally containing one or more Si or Ge atoms; more preferably, the substituents L are selected from the group consisting of -Cl, -Br, -Me, -Et, -n-Bu, -sec-Bu, -Ph, -Bz, -CH₂SiMe₃, -OEt, -OPr, -OBu, -OBz and -NMe₂.

The integer n ranges from 0 to 4, and it is preferably 1 or 2.

When $\mathbf{n} = 0$ and $\mathbf{r} = 1$, A can have only the meaning of Cp; Cp and A are preferably pentamethyl cyclopentadienyl, indenyl or 4,5,6,7-tetrahydroindenyl groups.

Non-limiting examples of these metallocene complexes are:

 $(Me_3Cp)_2MCl_2$ $(Me_4Cp)_2MCl_2$ $(Me_5Cp)_2MCl_2$ $(EtMe_4Cp)_2MCl_2$ $[(C_6H_5)Me_4Cp]_2MCl_2$ $(Et_5Cp)_2MCl_2$

 $(Ind)_3MCl_2$ $(H_4Ind)_2MCl_2$ $(Me_4Cp)(Me_5Cp)MCl_2$

 $[(Si(CH_3)_3Cp]_2MCl_2$ $(Me_5Cp)MCl_3$ $(Ind)MCl_3$ and

(H₄Ind)MCl₃

and the corresponding $-MMe_2$, $-M(OMe)_2$, $-M(OEt)_2$, -MCl(OMe), -MCl(OEt), $-MPh_2$, $-MBz_2$, -MMeCl, -MPhCl, $-M(NMe_2)_2$ and $-M(NMe_2)OMe$ derivatives, wherein Me = methyl, Et = ethyl, Cp = cyclopentadienyl, Ind = indenyl, $H_4Ind = 4,5,6,7$ -tetrahydroindenyl, Ph = phenyl, Ph

When n=1 or 2 and r=1, Cp and A, same or different from each other, are preferably cyclopentadienyl, tetramethyl-cyclopentadienyl, indenyl, 4,5,6,7-tetrahydroindenyl, 2-methyl-

4,5,6,7-tetrahydroindenyl, 4,7-dimethyl-4,5,6,7-tetrahydroindenyl, 2,4,7-trimethyl-4,5,6,7-tetrahydroindenyl or fluorenyl groups; $(ZR_m)_n$ is preferably Me₂Si, Me₂C, CH₂ or C₂H₄. Non-limiting examples of metallocene complexes of formula (I), wherein $\mathbf{n} = 1$ or 2 and $\mathbf{r} = 1$, are:

$Me_2Si(Me_4Cp)_2MCl_2$	Me ₂ C(Me ₄ Cp)(MeCp)MCl ₂	Me ₂ Si(Ind) ₂ MCl ₂
$C_2H_4(Ind)_2MCl_2$	$C_2H_4(H_4Ind)_2MCl_2$	Ph(Me)Si(Ind) ₂ MCl ₂
Ph ₂ Si(Ind) ₂ MCl ₂	Me ₂ C(Flu)(Cp)MCl ₂	•
$C_2H_4(Me_4Cp)_2MCl_2$	$C_2Me_4(Ind)_2MCl_2$	Me ₂ SiCH ₂ (Ind) ₂ MCl ₂
$C_2H_4(2-MeInd)_2MCl_2$	$C_2H_4(3-MeInd)_2MCl_2$	$C_2H_4(4,7-Me_2Ind)_2MCl_2$
$C_2H_4(5,6-Me_2Ind)_2MCl_2$	$C_2H_4(2\text{-MeH}_4\text{Ind})_2MCl_2$	$C_2H_4(2,4,7-Me_3H_4Ind)_2MCl_2$
$C_2H_4(4,7-Me_2H_4Ind)_2MCl_2$	$C_2H_4(2,4,7-Me_3Ind)_2MCl_2$	$C_2H_4(2\text{-Me-Benz}[e]Ind)_2MCl_2$
$C_2H_4(Benz[e]Ind)_2MCl_2$	Me ₂ Si(2-MeInd) ₂ MCl ₂	$Me_2Si(4,7-Me_2Ind)_2MCl_2$
Me ₂ Si(2-Me-4-Ph-Ind) ₂ MCl ₂	$Me_2Si(5,6-Me_2Ind)_2MCl_2$	Me ₂ Si(2,4,7-Me ₃ Ind) ₂ MCl ₂
Me ₂ Si(2-MeH ₄ Ind) ₂ MCl ₂	$Me_2Si(4,7-Me_2H_4Ind)_2MCl_2$	$Me_2Si(2,4,7-Me_3H_4Ind)_2MCl_2$
Me ₂ Si(Benz[e]Ind) ₂ MCl ₂	Me ₂ Si(2-Me-Benz[e]Ind) ₂ MCl	$_{2}$ Me $_{2}$ C(Ind) $_{2}$ MCl $_{2}$
$Me_2C(3-Me-Ind)_2MCl_2$	Me ₂ C(3-iPr-Ind) ₂ MCl ₂	Me ₂ C(3-Me ₃ Si-Ind) ₂ MCl ₂
$Me_2C(3-tBu-Ind)_2MCl_2$	$Me_2C(3-tBu-H_aInd)_2MCl_2$	$Me_2C(3-tBu-Cp)_2MCl_2$
$Me_2C(2-Me-4-tBu-Cp)_2MCl_2$	$H_2C(3-tBu-Ind)_2MCl_2$	H ₂ C(3-iPr-Ind) ₂ MCl ₂
H ₂ C(3-Me ₃ Si-Ind) ₂ MCl ₂	$H_2C(4,7-Me_2Ind)_2MCl_2$	$H_2C(1-Ph-5,7-Me_2Ind)_2MCl_2$
H ₂ C(2-Me-Ind) ₂ MCl ₂	H ₂ C(2-Me-3-Me ₃ Si-Ind) ₂ MCl ₂	$H_2C(Ind)_2MCl_2$
and the corresponding -MMe ₂	, -M(OMe) ₂ , -M(OEt) ₂ , -MCl(O	Me), -MCl(OEt), -MPh ₂ , -MBz ₂ , -
MMeCl, -MPhCl, -M(NMe ₂) ₂	and -M(NMe ₂)OMe derivative	es, wherein Me, Cp, Ind, Flu, Ph,
Bz, H ₄ Ind and M has the mean	ings reported above.	
Suitable metallocene complex	xes (A) are the bridged bis-inde	nyl metallocenes as described for

Suitable metallocene complexes (A) are the bridged bis-indenyl metallocenes as described for instance in USP 5,145,819 and EP-A-0 485 823.

Further metallocene complexes suitable for the catalyst system of the invention are the classes of heterocyclic metallocenes described in WO 98/22486 and WO 99/24446. Among these metallocenes, particularly preferred are the ones reported from page 15, line 8 to page 24, line 17; from page 25, line 1 to page 31, line 9; and from page 58, penultimate line, to page 63, line 20 of WO 98/22486. Other preferred metallocenes are the ones obtained from the bridged ligands listed from page 11, line 18, to page 14, line 13 of WO 99/24446

In the organometallic aluminium compounds (B) of formula (II), Ar is a substituted or

unsubstituted aryl group having from 6 to 20 carbon atoms; said aryl group has preferably formula (III):

$$R^{6}$$
 R^{7}
 R^{8}
(III)

wherein R^6 , R^8 and R^{10} are selected from the group consisting of hydrogen, halogen, $-R^3$, $-C(O)R^3$, $-OR^3$, $-SR^3$ $-NR^3$ ₂ and $-NO_2$, wherein R^3 has the meaning given above; and the groups R^7 and R^9 are selected from the group consisting of hydrogen, halogen, linear or branched, saturated or unsaturated C_1 - C_{20} alkyl, C_3 - C_{20} cycloalkyl, C_6 - C_{20} aryl, C_7 - C_{20} alkylaryl and C_7 - C_{20} arylalkyl groups, optionally containing one or more Si or Ge atoms; two adjacent substituents R^6 - R^{10} optionally form a ring, having 3 to 8 carbon atoms.

According to a particularly preferred embodiment of the invention, Ar is selected from the group consisting of 4-fluoro-phenyl, 4-chloro-phenyl, 4-methoxyphenyl, 4-nitrophenyl, 3-methylphenyl, 3-isopropylphenyl, 2,4-difluorophenyl, 2,4-dichlorophenyl, 2,6-difluorophenyl, 2,6-difluorophenyl, 3,5-difluorophenyl, 3,5-dichlorophenyl, 2,4,6-trifluorophenyl, 2,4,6-trifluorophenyl, 3,4,5-trifluorophenyl, 3,4,5-trichlorophenyl, pentafluorophenyl and pentachlorophenyl.

In the organometallic aluminium compound of formula (II), R^4 is preferably a C_1 - C_5 alkyl group, more preferably a C_1 - C_3 alkyl group, and even more preferably is methyl or ethyl; R^5 is preferably hydrogen or a C_1 - C_5 alkyl group, more preferably hydrogen or a C_1 - C_3 alkyl group, and even more preferably is hydrogen, methyl or ethyl; \mathbf{x} is 2 or 3, and preferably is 3.

According to the invention, component (B) can suitably comprise a mixture of two or more organometallic aluminium compounds of formula (II). Moreover, Component (B) can be used in combination with other organometallic aluminium compounds, different from those of formula (II), or in mixture with other compatible cocatalysts known in the state of the art.

In the organometallic aluminium compounds of formula (II), y is 0 or 1. As it is known in the state of the art, aluminium trialkyls may contain small amounts of bisalkyl-aluminium hydride, the hydride content can slightly change during prolonged storage periods and depending on the

storage temperature. Therefore, according to a preferred embodiment of the invention, component (B) is a mixture of the two organometallic aluminium compounds of formula (II) wherein y=0 and y=1; said mixtures of organometallic aluminium compounds may be prepared with methods known in the state of the art.

Non-limiting examples of organometallic aluminium compounds (B) of formula (II), according to the present invention, are:

tris(2-phenyl-propyl)aluminium,

tris[2-(4-fluoro-phenyl)-propyl]aluminium,

tris[2-(4-chloro-phenyl)-propyl]aluminium,

tris[2-(3-isopropyl-phenyl)-propyl]aluminium,

tris(2-phenyl-butyl)aluminium,

tris(3-methyl-2-phenyl-butyl)aluminium,

tris(2-phenyl-pentyl)aluminium,

tris[2-(pentafluorophenyl)-propyl]aluminium

tris[2,2-diphenyl-ethyl]aluminium

tris[2-phenyl-2-methyl-propyl]aluminium

as well as the corresponding compounds wherein one of the hydrocarbyl groups is replaced by a hydrogen atom. Particularly preferred compounds are tris(2-phenyl-propyl)aluminium, tris[2-(4-fluoro-phenyl)-propyl]aluminium and tris[2-(4-chloro-phenyl)-propyl]aluminium.

The components of the catalysts of the present invention can be brought into contact in different ways. It is possible for instance, to contact first the aluminium compound (B) with water (C) and subsequently to bring the thus obtained reaction product into contact with the metallocene complex (A).

Therefore, a preferred embodiment of the present invention is a catalyst system for the polymerization of olefins comprising the product obtained by contacting the following components:

(A) a metallocene complex of formula (I):

$$(Cp)(ZR_{m}^{1})_{n}(A)_{r}ML_{p} \qquad (I)$$

wherein M, Cp, $(ZR'_m)_p$, A, L, **r** and **p** have the meanings reported above; and

(B') the product of the reaction between water and an organometallic aluminium compound of formula (II):

Al
$$[CH_2-C(Ar)R^4R^5]_xH_y$$
 (II)

wherein Ar, R^4 , R^5 , x and y have the meanings reported above.

The molar ratio between said organometallic aluminium compound and said water preferably ranges from 1:1 to 100:1; more preferably from 1:1 to 50:1; even more preferably, the Al/H₂O molar ratio is 2. The expression "product of reaction" means the product obtained by contacting the above-mentioned components.

The molar ratio between said organometallic aluminium compound and said metallocene complex, expressed as Al/M molar ration, preferably ranges from 50 to 50000, more preferably from 500 to 5000.

A further object of the present invention is an alumoxane obtained by contacting an organometallic aluminium compound of formula (II):

Al
$$[CH_2-C(Ar)R^4R^5]_xH_y$$
 (II)

wherein Ar, R^4 , R^5 , x and y have the meanings reported above, with water, at a molar ratio between said organometallic aluminium compound and said water ranging from 1:1 to 100:1; preferably from 1:1 to 50:1, and even more preferably, at a Al/ H_2O molar ratio of about 2.

The compounds of formula (II), as reported above, may be prepared according to methods known in the state of the art. For instance, they may be prepared by reacting an alkene of formula $CH_2=C(Ar)R^4R^5$, wherein Ar, R^4 and R^5 have the meanings reported above, with an aluminium compound of formula AlR'3, wherein R' is hydrogen or a lower alkyl radical containing a β -hydrogen substituent; suitable AlR'3 compounds are triisobutylaluminium and diisobutylaluminium hydride.

The components (A), (B) and (C) of the catalysts of the present invention can be brought into contact by other methods known in the state of the art, such as by first contacting said organometallic aluminium compound with said metallocene complex and, thereafter, with water. According to an embodiment of the invention, water can be gradually added to said organometallic aluminium compound in solution, in an aliphatic or aromatic inert hydrocarbon solvent, such as heptane or toluene. Thereafter, the thus obtained solution is contacted with a solution of said metallocene complex in a suitable solvent, such as toluene.

According to another embodiment of the invention, water can be introduced in the monomer or in one of the monomers to be polymerized. In this case, said organometallic aluminium compound and said metallocene complex are pre-contacted before being used in the

polymerization.

According to another embodiment of the invention, water can be reacted in a combined form as a hydrated salt, or it can be adsorbed or absorbed on an inert support such as silica.

According to a further embodiment, said organometallic aluminium compound can be allowed to react with boric anhydride and with boric acid.

The catalyst system of the invention may be formed prior to its introduction into a polymerization reactor or *in situ* in the reactor, by contacting the above-described components.

According to a further embodiment, the catalyst system of the present invention is prepared according to the process described in the International Patent Application WO 99/21896, by prealkylating the metallocene complex (A) with an organometallic aluminium compound.

According to a preferred embodiment, the catalyst system of the invention is obtained by first contacting a metallocene complex of formula (I), as reported above, with one or more organometallic aluminium compounds of formula (IV):

$$AlR^{11}_{3-z}H_z$$
 (IV)

wherein R^{11} is a linear or branched, saturated or unsaturated, C_1 - C_{10} alkyl, C_6 - C_{20} aryl, C_7 - C_{20} arylalkyl or C_7 - C_{20} alkylaryl and z is 0 or 1; according to a preferred embodiment, said metallocene complex of formula (I) is first contacted with a mixture of two organometallic aluminium compounds of formula (IV) wherein z=0 and z=1, so that the molar ratio between the hydrogen atoms directly bound to aluminium and aluminium atoms is higher than 0.02. Examples of R^{11} substituents include 2,4,4-trimethyl-pentyl, 2,3,3-trimethyl-butyl and 2,3-dimethyl.

Subsequently the thus obtained mixture is contacted with (B'), i.e. the product of the reaction between an organometallic aluminium compound of formula (II), as reported above, and water.

The catalysts of the present invention can be used on inert supports. This may be achieved by depositing said metallocene complex, or the product of the reaction thereof with the aluminium compound pre-reacted with water, or said organometallic aluminium compound pre-reacted with water and subsequently said metallocene complex, on inert supports such as silica, alumina, styrene/divinylbenzene copolymers, polyethylene or polypropylene.

The thus obtained solid compound, together with further addition of said organometallic aluminium compound, either as such or pre-reacted with water, can be suitably used in gas phase polymerization.

The catalysts of the present invention can be used in the polymerization reactions of olefins.

Therefore, according to further object, the invention provides a process for the polymerization of one or more olefins in the presence of a catalyst system as described above. Olefins which can be polymerized with the process of the present invention are, for instance, α -olefins of formula CH,=CHR, wherein R is hydrogen or a C_1 - C_{20} alkyl radical.

The catalysts according to the present invention can be conveniently used in the homopolymerization of ethylene, in particular for the preparation of HDPE, and in the copolymerization of ethylene, in particular for the preparation of LLDPE. Suitable comonomers in ethylene copolymers are α -olefins of formula CH₂=CHR', wherein R' is a linear, branched or cyclic C_1 - C_{20} alkyl radical, and cycloolefins. Examples of such olefins are propylene, 1-butene, 1-pentene, 4-methyl-1-pentene, 1-hexene, 1-octene, allyl-cyclohexane, cyclopentene, cyclohexene, norbornene and 4,6-dimethyl-1-heptene.

Further suitable comonomers in said ethylene copolymers are polyenes, in particular conjugated or non-conjugated, linear or cyclic dienes, such as 1,4-hexadiene, isoprene, 1,3-butadiene, 1,5-hexadiene and 1,6-heptadiene.

The catalysts of the invention can be suitably used in propylene homopolymerization, in particular for the production of isotactic polypropylene.

Moreover, the catalysts of the invention can be suitably used in the preparation of elastomeric copolymers of ethylene with α -olefins of formula CH₂=CHR, wherein R is a C₁-C₁₀ alkyl radical, such as propylene, 1-butene, 4-methyl-1-pentene, 1-hexene and 1-octene.; said copolymers may optionally contain minor proportions of units deriving from polyenes.

According to a further embodiment, the catalysts according to the present invention are used in the preparation of cycloolefin polymers. Monocyclic and polycyclic olefin monomers can be either homopolymerized or copolymerized, also with linear olefin monomers. Non limiting examples of cycloolefin polymers which can be prepared with the catalyst of the present invention are described in the European patent applications EP 0 501 370 and EP 0 407 870.

The polymerization processes of the present invention can be carried out in liquid phase, optionally in the presence of an inert hydrocarbon solvent, or in gas phase. Said hydrocarbon solvent can be either aromatic (such as toluene) or aliphatic (such as propane, hexane, heptane, isobutane, cyclohexane and 2,2,4-trimethylpentane).

The polymerization temperature preferably ranges from 0°C to 250°C; in the preparation of

PCT/EP00/09111

WO 01/21674

HDPE and LLDPE, it is preferably comprised between 20°C and 150°C and, more particularly between 40°C and 90°C; in the preparation of elastomeric copolymers, it is preferably comprised between 0°C and 200°C, and more preferably between 20°C and 100°C. The catalyst systems according to the invention have the advantage of maintaining very high activity also when relatively high polymerization temperatures are used.

The molecular weight of the (co)polymers can be varied simply by varying the polymerization temperature, the type or the concentration of the catalyst components, or by using molecular weight regulators, such as hydrogen.

The molecular weight distribution can be varied by using mixtures of different metallocene complexes or by carrying out the polymerization in several stages which differ in the polymerization temperature and/or the concentrations of molecular weight regulator.

The polymerization yield depends on the purity of the metallocene complex (A) in the catalyst, therefore, said metallocene complex can be used as such or can be subjected to purification treatments before use.

Particularly interesting results are obtained when the components of the catalyst of the invention are contacted among them before the polymerization. The contact time is preferably comprised between 1 and 60 minutes, more preferably between 5 and 20 minutes. The precontact concentrations for the metallocene complex are comprised between 10^{-2} and 10^{-8} mol/l, while for the product of the reaction between the organometallic aluminium compound and water they are comprised between 10 and 10^{-3} mol/l. The precontact is preferably carried out in the presence of a hydrocarbon solvent and, optionally, of small amounts of monomer.

The following examples are given for illustrative and not limiting purposes.

SYNTHESIS OF METALLOCENE COMPLEXES

rac-Ethylene-bis(1-indenyl)zirconium dichloride

rac-Et(1-Ind)₂ZrCl₂

The metallocene complex was prepared according to the procedure described in EP-A-575,875.

rac-Dimethylsilyl-bis(2-methyl-1-indenyl)zirconium dichloride rac-Me,Si(2-Me-1-Ind),ZrCl₂

The metallocene complex was obtained from Boulder Scientific Company as the 97 % rac isomer and was used without further purification.

rac-Methylene-bis(3-t-butyl-1-indenyl)zirconium dichloride

rac-CH₂(3-tBu-1-Ind)₂ZrCl₂

The metallocene complex was prepared as described in Synthesis 1 of the International Patent Application WO 98/43989.

rac-Dimethylsilyl-bis(2-methyl-benzoindenyl)zirconium dichloride

Me,Si(2-Me-Benzind),ZrCl,

The metallocene complex was prepared as described in Example A of the U.S. Patent 5,455,366.

SYNTHESIS OF ORGANOALUMINIUM COMPOUNDS

General Procedure

All reactions were carried out under nitrogen in the glove box or under Schlenk conditions using oven-dried glassware. The toluene solvent was dried over 4Å molecular sieves. All alkenes were dried over 4Å molecular sieves prior to use.

Tris(2-phenyl-propyl)aluminium - Al(CH2CHMePh)3

In a glove box, o-methyl-styrene (283g, 2.3 mol; Aldrich, dried over sieves) was dissolved in dry toluene (ca. 300 ml) in a 1 L 3-neck flask. Al{CH₂CHMe₂}₃ (TIBA, 100 ml, 0.395 mmol, ex-Witco) was added over 10 min by syringe to the rapidly stirred solution at ambient temperature. The reaction flask was removed from the glove box and a reflux condenser and nitrogen line attached in the fume hood. The isobutene product was collected using a graduated collection vessel immersed in a -78°C acetone/dry ice bath. The reaction mixture was warmed over 90 minutes to an internal temperature of 110.7 °C. The reaction was allowed to reflux for 16 hours (final reflux temperature 126.4 °C), affording ca. 100% of the theoretical maximum yield of isobutene (ca. 3.0 equivalents/Al). The remaining olefin and solvent were removed *in vacuo* (50°C, 0.05 mbar, 90 min) utilizing a dry ice/acetone bath to give 162 g of tris(2-phenyl-propyl)aluminium.

Tris[2-(4-fluoro-phenyl)-propyl]aluminium - Al[CH₂CHMe(4-F-C₆H₄)]₃

In the glove box, 2-(4-fluoro-phenyl)-propylene (65.1 g, 0.48 mol; Acros, dried over sieves) was dissolved in dry toluene (ca. 70 ml) in a 250 ml 3-neck flask. Al{CH₂CHMe₂}₃ (TIBA, 27.9 ml, 0.120 mol, ex-Witco) was added over 10 min by syringe to the rapidly stirred solution. The reaction flask was removed from the glove box and a reflux condenser and nitrogen line attached in the fume hood. The isobutene product was collected using a

PCT/EP00/09111

WO 01/21674

graduated collection vessel immersed in a -78°C acetone/dry ice bath. The reaction mixture was warmed over 90 minutes to an internal temperature of 119.6 °C. The reaction was allowed to reflux for 16 hours (final reflux temperature 123.5°C), affording ca. 100 % of the theoretical maximum yield of isobutene. The remaining olefin and solvent were removed *in vacuo* (60°C, 0.05 mbar, 90 min) utilizing a dry ice/acetone bath to give 50 g of tris[2-(4-fluoro-phenyl)-propyl]aluminium.

$Tris[2\hbox{-}(4\hbox{-}chloro\hbox{-}phenyl)\hbox{-}propyl] a luminium - Al[CH_2CHMe(4\hbox{-}Cl\hbox{-}C_6H_4)]_3$

In the glove box, 2-(4-chloro-phenyl)-propylene (73.2 g, 0.48 mol; Acros, dried over sieves) was dissolved in dry toluene (ca. 80 ml) in a 250 ml 3-neck flask. Al{CH₂CHMe₂}₃ (TIBA, 30.0 ml, 0.128 mol, ex-Witco) was added over 10 min by syringe to the rapidly stirred solution. The reaction flask was removed from the glove box and a reflux condenser and nitrogen line attached in the fume hood. The isobutene product was collected using a graduated collection vessel immersed in a -78°C acetone/dry ice bath. The reaction mixture was warmed over 90 minutes to an internal temperature of 123.4 °C. The reaction was allowed to reflux for 18 hours (final reflux temperature 124.4 °C), affording ca. 100 % of the theoretical maximum yield of isobutene. The remaining olefin and solvent were removed *in vacuo* (60°C, 0.05 mbar, 90 min) utilizing a dry ice/acetone bath to give 62 g of tris[2-(4-chloro-phenyl)-propyl]aluminium.

$Tris[2\hbox{-}(4\hbox{-methyl-phenyl})\hbox{-propyl}] a luminium - Al[CH_2CHMe(4\hbox{-Me-C}_6H_4)]_3$

In the glove box, 2-(4-methyl-phenyl)-propylene (39.5 g, 0.299 mol) was dissolved in dry toluene (ca. 45 ml) in a 250 ml 3-neck flask. Al{CH₂CHMe₂}₃ (TIBA, 18.7 ml, 0.080 mol, ex-Witco; Aldrich, dried over sieves) was added over 10 min by syringe to the rapidly stirred solution. The reaction flask was removed from the glove box and a reflux condenser and nitrogen line attached in the fume hood. The isobutene product was collected using a graduated collection vessel immersed in a -78°C acetone/dry ice bath. The reaction mixture was warmed over 60 minutes to an internal temperature of 124.3 °C. The reaction was allowed to reflux for 20 h (final reflux temperature 125.4 °C), affording ca. 100 % of the theoretical maximum yield of isobutene. The remaining olefin and solvent were removed *in vacuo* (60 °C, 0.05 mbar, 90 min) utilizing a dry ice/acetone bath to give 30 g of tris[2-(4-methyl-phenyl)-propyl]aluminium.

 $Tris[2-(4-isopropyl-phenyl)-propyl] a luminium - Al[CH_2CHMe(4-iPr-C_6H_4)]_3$

In the glove box, 2-(4-isopropyl-phenyl)-propylene (27.5 g, 0.172 mol) was dissolved in dry toluene (ca. 35 ml) in a 500 ml 3-neck flask. Al{CH₂CHMe₂}₃ (TIBA, 10.8 ml, 0.046 mol, ex-Witco) was added over 10 min by syringe to the rapidly stirred solution. The reaction flask was removed from the glove box and a reflux condenser and nitrogen line attached in the fume hood. The isobutene product was collected using a graduated collection vessel immersed in a -78°C acetone/dry ice bath. The reaction mixture was warmed over 90 minutes to an internal temperature of 125.6 °C. The reaction was allowed to reflux for 20 h (final reflux temperature 131.8 °C), affording ca. 100 % of the theoretical maximum yield of isobutene. The remaining olefin and solvent were removed *in vacuo* (75 °C, 0.05 mbar, 90 min) utilizing a dry ice/acetone bath to give 25 g of tris[2-(4-isopropyl-phenyl)-propyl]aluminium.

Tris[2-(3-isopropyl-phenyl)-propyl] aluminium $-Al[CH_2CHMe(3-iPr-C_6H_4)]_3$

In the glove box, 2-(3-isopropyl-phenyl)-propylene (43.2 g, 0.27 mol; Chemsampco, dried over sieves) was dissolved in dry toluene (ca. 50 ml) in a 250 ml 3-neck flask. Al{CH₂CHMe₂}₃ (TIBA, 17.0 ml, 0.068 mol, ex-Witco) was added over 10 min by syringe to the rapidly stirred solution. The reaction flask was removed from the glove box and a reflux condenser and nitrogen line attached in the fume hood. The isobutene product was collected using a graduated collection vessel immersed in a -78°C acetone/dry ice bath. The reaction mixture was warmed over 60 minutes to an internal temperature of 124.1 °C. The reaction was allowed to reflux for 16 h (final reflux temperature 125.4 °C), affording ca. 100 % of the theoretical maximum yield of isobutene. Despite the low boiling point of CH₂=C(Me)(CMe₃), only a negligible amount of this olefin was collected in the cold trap. The remaining olefin and solvent were removed *in vacuo* 75 °C, 0.05 mbar, 90 min) utilizing a dry ice/acetone bath to give 32 g of tris[2-(3-isopropyl-phenyl)-propyl]aluminium.

Tris(2,3-dimethyl-butyl)aluminium

The aluminium compound was prepared as described in the International Patent Application WO 99/21899.

Tris(2,4,4-trimethyl-pentyl)aluminium (TIOA)

The aluminium compound was prepared according to the method described in Liebigs Ann. Chem., Volume 629, 1960, Ziegler et al. "Aluminiumtrialkyle und Dialkyl-aluminiumhydride aus Aluminiumisobutyl-Verbindungen [Aluminium trialkyls and dialkyl-aluminium hydrides from aluminium isobutyl compounds]", pages 14-19.

PCT/EP00/09111

Tris[2-(2-naphthyl)-propyl]aluminium - AI{CH2CHMe(Naphthyl)}3

In a glove box, 2-isopropenylnaphthalene (20.3 g. 121 mmol; TCI-US, evacuated for 2 h at 50 °C) was dissolved in dry toluene (ca. 30 ml) in a 250 mL 3-neck flask. Al{CH2CHMe2}3 (TIBA, 7.23 g, 36.5 mmol, AKZO) was added by syringe to the rapidly stirred solution at ambient temperature. The reaction flask was removed from the glove box and a reflux condenser and nitrogen line attached in the fume hood. The isobutene product was collected using a graduated collection vessel immersed in a -78°C acetone/dry ice bath. The reaction mixture was warmed over 20 minutes to an internal temperature of 120 °C. The reaction was allowed to reflux for 16 hours (final reflux temperature 120 °C), affording ca. 100% of the theoretical maximum yield of isobutene (ca. 3.0 equivalents/Al). The solvent was removed in vacuo (60°C, 0.03 mbar, 120 min) utilizing a dry ice/acetone bath to give 19.3 g of tris[2-(2-naphthyl)-propyl]aluminium, containing ca. 10 w/w % 2-isopropenylnaphthalene, as determined by 1H NMR spectroscopy.

Methylalumoxane (MAO)

The product available from Witco as a 10% w/w solution in toluene was used.

Tris(2-methyl-propyl)aluminium (TIBA)

The product available from Witco was used.

POLYMERIZATION

EXAMPLES 1-3

WO 01/21674

A 5 liters reactor equipped with turbine stirrer, steam/water temperature control and a catalyst injection system was heated to 150-160°C overnight, whilst purging with nitrogen, cooled and then pickled at 70°C using a mixture of TIBA (0.25 g), toluene (20 ml) and propylene (500 g). The pickle mixture was removed and the reactor then charged with 1650 g liquid propylene, whilst increasing the temperature from 20°C to 50°C.

Separately, 3.46 g of Tris(2-phenyl-propyl)aluminium Al(CH₂CHMePh)₃ (9 mmol) were dissolved in 20 g of toluene in a bottle with a septum cap. The solution was cooled to 0-4 °C using an ice bath, and 81 microliter of water (4.5 mmol) added in four shots using a 25 microliter syringe, whilst maintaining the temperature below 15°C and purging the solution with nitrogen The resulting solution was introduced into the reactor using an injection system, washed in using 20 ml of toluene.

Meanwhile, 10.1 mg of rac-Et(1-Ind)₂ZrCl₂ (24 micromol) was dissolved in 15.7 g of toluene,

and 0.65 g of the obtained solution was reacted with 0.11 g TIOA (0.3 mmol), resulting in a color change from yellow to light yellow. Ten minutes after the introduction of the hydrolyzed alkylaluminium mixture into the reactor, the alkylated zirconocene solution (aged for 5 minutes) was injected into the reactor (using 20 ml toluene). The polymerization was continued for 1 hour, at the temperature reported in table 1, using 840-1100 rpm stirring; in Example 3, moreover, hydrogen was added during the polymerization (0.05 %v of the gascap). The polymerization was then stopped by injection of 5-10 ml methanol. The heating was then discontinued and the propylene rapidly vented and the powder polypropylene collected. Fouled material was removed using hot toluene and precipitated with methanol. The polypropylene fractions were dried (70-80°C, 200 mbar, nitrogen purge) and combined to give the total yield of polypropylene.

The polymerization conditions and the data relating to the obtained polymers are indicated in Table 1.

COMPARATIVE EXAMPLES 1 AND 2

The general procedure described in Example 1 was followed, but using MAO instead of Al(CH₂CHMePh)₃ and instead of TIOA, and operating with the amounts of metallocene and aluminium compound indicated in Table 1; in comparative Example 2, moreover, hydrogen was added during the polymerization (0.05 %v of the gascap); the polymerization conditions and the data relating to the obtained polymer are indicated in Table 1.

COMPARATIVE EXAMPLES 3 AND 4

The general procedure described in Example 1 was followed, but using tris(2,3-dimethylbutyl)aluminium instead of Al(CH₂CHMePh)₃ and instead of TIOA, and using the amounts of metallocene and aluminium compound indicated in Table 1 (Al/H₂O molar ratio 2/1 during alumoxane preparation); the polymerization was carried out at 50°C and 70°C respectively. The polymerization conditions and the data relating to the obtained polymer are indicated in Table 1.

EXAMPLES 4 AND 5

The general procedure described in Example 1 was followed, but using Al[CH₂CHMe(4-F- C_6H_4)]₃ as aluminium compound instead of Al(CH₂CHMePh)₃ and using the amounts of metallocene and aluminium compound indicated in Table 1 (Al/H₂O molar ratio 2/1 during alumoxane preparation). The polymerization conditions and the data relating to the obtained

polymers are indicated in Table 1.

EXAMPLE 6

The general procedure described in Example 1 was followed, but using Al[CH₂CHMe(4-Cl- C_6H_4)]₃ as aluminium compound instead of Al(CH₂CHMePh)₃ and using the amounts of metallocene and aluminium compound indicated in Table 1 (Al/H₂O molar ratio 2/1 during alumoxane preparation). The polymerization conditions and the data relating to the obtained polymer are indicated in Table 1.

EXAMPLE 7

A 5 liters reactor equipped with turbine stirrer, steam/water temperature control and a catalyst injection system was heated to 150-160°C overnight, whilst purging with nitrogen, cooled and then pickled at 70°C using a mixture of TIBA (0.25 g), toluene (20 ml) and propylene (500 g). The pickle mixture was removed and the reactor then charged with 1650 g liquid propylene, whilst increasing the temperature from 20°C to 50°C.

Separately, 1.97 g of Al[CH₂CHMe(4-F-C₆H₄)]₃ (4.5 mmol) were dissolved in 20 g of toluene in a bottle with a septum cap. The solution was cooled to 0-4°C using an ice bath, and 41 microliter of water (2.25 mmol) added in four shots using a 25 microliter syringe, whilst maintaining the temperature below 15°C and purging the solution with nitrogen. The resulting solution was introduced into the reactor using an injection system, washed in using 20 ml of toluene.

Meanwhile, 9.2 mg of *rac*-CH₂(3-*t*Bu-1-Ind)₂ZrCl₂ (17.8 micromol) were dissolved in 15.8 g of toluene, and 2.67 g of the obtained solution was reacted with 0.11 g TIOA (0.3 mmol), resulting in a color change from red to pale red. Ten minutes after the introduction of the hydrolyzed alkylaluminium mixture into the reactor, the alkylated zirconocene solution (aged for 5 minutes) was injected into the reactor (using 20 ml toluene). The polymerization was continued for 1 hour, at the temperature of 70°C, using 840-1100 rpm stirring; hydrogen was added during the polymerization (0.07 %v of the gascap). The polymerization was then stopped by injection of 5-10 ml methanol. The heating was then discontinued and the propylene rapidly vented and the powder polypropylene collected. Fouled material was removed using hot toluene and precipitated with methanol. The polypropylene fractions were dried (70-80°C, 200 mbar, nitrogen purge) and combined to give the total yield of polypropylene.

The polymerization conditions and the data relating to the obtained polymer are indicated in Table 1.

EXAMPLE 8

A 5 liters reactor equipped with turbine stirrer, steam/water temperature control and a catalyst injection system was heated to 150-160°C overnight, whilst purging with nitrogen, cooled and then pickled at 70°C using a mixture of TIBA (0.25 g), toluene (20 ml) and propylene (500 g). The pickle mixture was removed and the reactor then charged with 1650 g liquid propylene, whilst increasing the temperature from 20°C to 50°C.

Separately, 4.37 g of Al[CH₂CHMe(4-F-C₆H₄)]₃ (10 mmol) was dissolved in 20 g of toluene in a bottle with a septum cap. The solution was cooled to 0-4 °C using an ice bath, and 81 microliter of water (4.5 mmol) added in four shots using a 25 microliter syringe, whilst maintaining the temperature below 15°C and purging the solution with nitrogen. The resulting solution was introduced into the reactor using an injection system, washed in using 20 ml of toluene.

Meanwhile, 10.3 mg of rac-Me₂Si(2-Me-1-Ind)₂ZrCl₂ (21.7 micromol) was dissolved in 15.6 g of toluene, and 0.72 g of the obtained solution was reacted with 0.11 g TIOA (0.3 mmol), resulting in a color change from yellow to light yellow. Ten minutes after the introduction of the hydrolyzed alkylaluminium mixture into the reactor, the alkylated zirconocene solution (aged for 5 minutes) was injected into the reactor (using 20 ml toluene). The polymerization was continued for the time reported in Table 1, at the temperature reported in Table 1, using 840-1100 rpm stirring. The polymerization was then stopped by injection of 5-10 ml methanol.

The polymerization conditions and the data relating to the obtained polymers are indicated in Table 1.

EXAMPLE 9

The general procedure described in Example 8 was followed, using the amounts of metallocene and aluminium compound indicated in Table 1 (Al/H₂O molar ratio 2/1 during alumoxane preparation); hydrogen was added during the polymerization (0.24 %v of the gascap). The polymerization conditions and the data relating to the obtained polymers are indicated in Table

EXAMPLES 10-12

The general procedure described in Example 8 was followed, but using Al[CH₂CHMe(4-Cl- C_6H_4)]₃, as aluminium compound instead of Al[CH₂CHMe(4-F- C_6H_4)]₃, and using the amounts of metallocene and aluminium compound indicated in Table 1 (Al/H₂O molar ratio 2/1 during alumoxane preparation); in Example 12, moreover, hydrogen was added during the polymerization (0.11 %v of the gascap). The polymerization conditions and the data relating to the obtained polymers are indicated in Table 1.

COMPARATIVE EXAMPLE 5

The general procedure described in Example 8 was followed, but using MAO as aluminium compound instead of $Al[CH_2CHMe(4-F-C_6H_4)]_3$, and using the amounts of metallocene, and aluminium compound indicated in Table 1; hydrogen was added during the polymerization (0.11 %v of the gascap). The polymerization conditions and the data relating to the obtained polymers are indicated in Table 1.

EXAMPLE 13

A 5 liters reactor equipped with turbine stirrer, steam/water temperature control and a catalyst injection system was heated to 150-160°C overnight, whilst purging with nitrogen, cooled and then pickled at 70°C using a mixture of TIBA (0.25 g), toluene (20 ml) and propylene (500 g). The pickle mixture was removed and the reactor then charged with 1650 g liquid propylene, whilst increasing the temperature from 20°C to 30°C. Then 4-5 % hydrogen was added to the gascap, aiming at 1-1.5% hydrogen in the gascap at 70°C.

Separately, 2.20 g of tris {2-(4-chlororophenyl)-propyl} Al[CH₂CHMe(4-Cl-C₆H₄)]₃ (4.5 mmol) were dissolved in 20 ml of toluene in a bottle with a septum cap. The solution was cooled to 0-4 °C using an ice bath, and 41 microliter of water (2.28 mmol) added in two shots using a 100 microliter syringe, whilst maintaining the temperature below 15°C. The resulting solution was introduced into the reactor using an injection system, washed in using 20 ml of toluene.

Meanwhile, 13 mg of rac-Me₂Si(2-Me-Benzind)₂ZrCl₂ (23 micromol) was dissolved in 22.39 g of toluene by adding 413 mg TIOA (1128 micromol), and 0.05 g of the obtained solution containg 0.05 micromol zirconium complex was reacted with 0.11 g TIOA (0.3 mmol), resulting in a color change from yellow to light yellow. Ten minutes after the introduction of the hydrolyzed alkylaluminium mixture containing 4.5 mmol aluminoxane into the reactor, the alkylated zirconocene solution (aged for 5 minutes) was injected into the reactor (using 20 ml toluene) at a temperature of 30°C. After 0.5 minute the temperature was raised in 6-7 minutes to

70°C and polymerization was continued for 1 hour, using 840-1100 rpm stirring, keeping the hydrogen concentration at 1 % in the gascap. The polymerization was then stopped by injection of 5-10 ml methanol. The heating was then discontinued and the propylene rapidly vented and the powder polypropylene collected. Fouled material was removed using hot xylene and precipitated with methanol. The polypropylene fractions were dried (70-80°C, 200 mbar, nitrogen purge) and combined to give the total yield of polypropylene.

COMPARATIVE EXAMPLE 6

A 5 liters reactor equipped with turbine stirrer, steam/water temperature control and a catalyst injection system was heated to 150-160°C overnight, whilst purging with nitrogen, cooled and then pickled at 70°C using a mixture of TIBA (0.25 g), toluene (20 ml) and propylene (500 g). The pickle mixture was removed and the reactor then charged with 1650 g liquid propylene, whilst increasing the temperature from 20°C to 30°C. Then 4-5 % hydrogen was added to the gascap, aiming at 1-1.5% hydrogen in the gascap at 70°C.

Separately, 2.44 g of a toluene solution of MAO containing 4.98 % w/w aluminium (4.5 mmol) was introduced into the reactor using an injection system, washed in using 20 ml of toluene..

Meanwhile, 17.1 mg of Me₂Sì(2-Me-Benzind)₂ZrCl₂ (30 micromol) was dissolved in 16.46 g of toluene by adding 138 mg MAO solution (250 micromol), and 0.745 g of the obtained solution containing 0.05 micromol zirconium complex was reacted with 405 mg of the MAO solution (0.75 mmol), resulting in a color change from yellow to red. Ten minutes after the introduction of the solution containing 4.5 mmol MAO into the reactor, the alkylated zirconocene solution (aged for 5 minutes) was injected into the reactor (using 20 ml toluene) at a temperature of 30°C. After 0.5 minute the temperature was raised in 6-7 minutes to 70°C and polymerization was continued for 1 hour, using 840-1100 rpm stirring, keeping the hydrogen concentration at 1 % in the gascap. The polymerization was then stopped by injection of 5-10 ml methanol. The heating was then discontinued and the propylene rapidly vented and the powder polypropylene collected. Fouled material was removed using hot xylene and precipitated with methanol. The polypropylene fractions were dried (70-80°C, 200 mbar, nitrogen purge) and combined to give the total yield of polypropylene.

		(longing)	Orgality ating agent	Cocatalyst (mmol	(mmol) Total	် (၁ (၁		ıme	Activity
Example	Metallocene	(micronioi)	(Jones)		Al/Zr		(g pol.)	(min)	(g pol.) (min) (kg/g Zr/h)
*			(ionum)		(mol/mol)				
	100		FIO.	AI/CH2CHMePh)3 9	9300	20	615	31	3530
-	rac-Et(1-Ind)2ZrCl2			C/	0300	70	450	09	4930
2	rac-Et(1-Ind)2ZrCl2	1	TIOA 0.3		+	2 5	204	61	6480
3*	rac-Et(1-Ind)2ZrCl2	1	TIOA 0.3	AI(CH2CHMePh)3	\dashv	2 5	- 00	5 6	0000
Comp 1	Comp 1 rac-Et(1-Ind)2CClo	-	MAO 0.5	MAO 8.8	-	2	382	3 3	0004
to deligo	2 2	-	MAO 0.5	MAO	9 9540	70	295	2.1	0000
Comp.z	190-EU 1-1110/251012	6	AI/CH2CHMeCHMe2)3 0.3	AI(CH2CHMeCHMe2)3	9 4650	20	432	09	2470
Comp.3	Comp.3 rac-Et(1-1110)221C12	4 0		AIICH2CHMeCHMe2)3	9 4650	70	152	09	830
Comp.4	Comp.4 rac-Et(1-Ind)22rCl2	7	١,	135	9 18400	20	395	09	8660
4	rac-Et(1-Ind)2ZrCl2	0.5	IIOA U.2	l	1	5	455	61	9810
2	rac-Et(1-Ind)2ZrCl2	0.5	TIOA 0.2	١	00400	2 6	700	. 0	6290
9	rac-Et(1-Ind)2ZrCl2	0.5	TIOA 0.3	AI[CH2CHMe(4-CI-C6H4)]3	184/0	2	107	3	
		١		3 VICUSCHMS(4 E-CCH4)19 45	1600	2	160	09	580
7*	Jac-CH2(3-tBu-1-Ind)2ZrCl2	·Cl2 3	IIOA U.3	1	+	70	135**	10	8880**
α	rac-MeoSi(2-Me-1-Ind)oZrClo	ZrClo 1	TIOA 0.3	AICH2CHMe(4-F-C6H4)]3 10	4	2	3	2 :	0232
>	CIDZC(pal-1-aW C):2coW con	7rClo 0.5	TIOA 0.3	AI[CH2CHMe(4-F-C6H4)]3	2 4600	20	239	4	0/0/
5	180-1916201(2-1916-1-1916)2-1012	- 1	TIOA 0.3	AICH2CHMe(4-CI-C6H4)]3 4.5	5 19200	20	113	99	4960
2	/ac-ivie25i(z-ivie-1-iiiu)	- 1		AIICH2CHMe(4-CI-C6H4)]3 4.5	5 19200	2	283	99	12400
-	rac-Me ₂ Si(Z-Me-1-Ind) ₂ ZrCi ₂	- 1		1	2 4600	2	236.	ල _	10350**
12*	rac-Me ₂ Si(2-Me-1-Ind) ₂ ZrCl ₂	ι		Cut. 100 10 1 107	+	2	62	25	4160
Comp.5*	Comp.5* rac-Me2Si(2-Me-1-Ind)2ZrCl2	ZrCl ₂ 0.5	TIOA 0.3	- 1	-	2 6		e G	34900
13	rac-Me2Si(2-Me-Benzind)2ZrC	d)2ZrCl2 0.05	TIOA 0.3	Al[CH2CHMe(4-CI-C6H4)]3 4.5	\dashv	2		3 8	20200
Comp	Comp. 6 rac-Me2Si/2-Me-Benzind)2ZrC	d)2ZrCl2 0.05	MAO 0.75	MAO	4.5 90010	2		20	20.00
7	ומכיוויים בייים						•		

Hydrogen added during polymerization
 Calculated yield and activity (based on propene conversion)

CLAIMS

- 1. A catalyst system for the polymerization of olefins comprising the product obtained by contacting the following components:
 - (A) a metallocene complex of formula (I):

$$(Cp)(ZR_{m}^{1})_{n}(A)_{n}ML_{p}$$
 (I)

wherein $(ZR_{m}^{1})_{n}$ is a divalent group bridging Cp and A, Z being C, Si, Ge, N or P, and the R¹ groups, equal or different from each other, being hydrogen or linear or branched, saturated or unsaturated C_{1} - C_{20} alkyl, C_{3} - C_{20} cycloalkyl, C_{6} - C_{20} aryl, C_{7} - C_{20} alkylaryl or C_{7} - C_{20} arylalkyl groups;

Cp is a substituted or unsubstituted cyclopentadienyl group, optionally condensed to one or more substituted or unsubstituted, saturated, unsaturated or aromatic rings, containing from 4 to 6 carbon atoms, optionally containing one or more heteroatoms;

A is -O-, -S-, -N(\mathbb{R}^2)-, wherein \mathbb{R}^2 is hydrogen, a linear or branched, saturated or unsaturated C_1 - C_{20} alkyl, C_3 - C_{20} cycloalkyl, C_6 - C_{20} aryl, C_7 - C_{20} alkylaryl or C_7 - C_{20} arylalkyl, or A has the same meaning of $\mathbb{C}p$;

M is a transition metal belonging to group 3, 4, 5, 6 or to the lanthanide or actinide groups of the Periodic Table;

the substituents L, same or different from each other, are monoanionic sigma ligands selected from the group consisting of hydrogen, halogen, $-R^3$, $-OR^3$, $-OCOR^3$, $-SR^3$, $-NR^3$ ₂ and $-PR^3$ ₂, wherein R^3 is a linear or branched, saturated or unsaturated C_1 - C_{20} alkyl, C_3 - C_{20} cycloalkyl, C_6 - C_{20} aryl, C_7 - C_{20} alkylaryl or C_7 - C_{20} arylalkyl group, optionally containing one or more Si or Ge atoms;

m is 1 or 2; n is an integer ranging from 0 to 4; r is 0 or 1; n is 0 when r is 0;

p is an integer equal to the oxidation state of the metal M minus 2 when r=1, and minus 1 when r=0, and ranges from 1 to 4;

(B) an organometallic aluminium compound of formula (II):

Al
$$[CH_2-C(Ar)R^4R^5]_xH_y$$
 (II)

wherein Ar is a substituted or unsubstituted aryl group having from 6 to 20 carbon atoms;

 R^4 is a linear or branched, saturated or unsaturated, C_1 - C_{10} alkyl, C_6 - C_{20} aryl, C_7 - C_{20} arylalkyl or C_7 - C_{20} alkylaryl; R^5 is hydrogen or a linear or branched, saturated or

unsaturated, C_1 - C_{10} alkyl, C_6 - C_{20} aryl, C_7 - C_{20} arylalkyl or C_7 - C_{20} alkylaryl group; the substituents R^4 and R^5 optionally form a ring, having 3 to 8 carbon atoms; a carbon atom in the compound of formula (II) being optionally replaced by a Si or a Ge atom;

x is 2 or 3; y = 3-x; and

- (C) water;
- the molar ratio between said organometallic aluminium compound (B) and said water (C) being comprised between 1:1 and 100:1.
- 2. The catalyst system according to claim 1, wherein the molar ratio between said organometallic aluminium compound (B) and said water (C) is about 2.
- 3. The catalyst system according to claim 1 or 2, wherein the molar ratio between said organometallic aluminium compound (B) and said metallocene complex (A) ranges from 50 to 50,000.
- 4. The catalyst system according to claim 1 wherein, in said metallocene complex of formula (I), M is Ti, Zr or Hf.
- 5. The catalyst system according to claim 1 wherein, in said metallocene complex of formula (I), the divalent bridge $(ZR_m^1)_n$ is selected from the group consisting of CR_2^1 , $(CR_2^1)_2$, $(CR_2^1)_3$, SiR_2^1 , GeR_2^1 , NR_2^1 and PR_2^1 , R_2^1 having the meaning reported in claim 1.
- 6. The catalyst system according to claim 5, wherein said divalent bridge $(ZR_m^1)_n$ is selected from the group consisting of $Si(CH_3)_2$, $SiPh_2$, CH_2 , $(CH_2)_2$, $(CH_2)_3$ and $C(CH_3)_2$.
- 7. The catalyst system according to claim 1 wherein, in said metallocene complex of formula (I), Cp is selected from the group consisting of cyclopentadienyl, mono-, di-, tri- and tetra-methyl cyclopentadienyl; 4-'butyl-cyclopentadienyl; 4-adamantyl-cyclopentadienyl; indenyl; mono-, di-, tri- and tetra-methyl indenyl; 3-'butyl-indenyl; 3-trimethylsilyl-indenyl; 4,5,6,7-tetrahydroindenyl; fluorenyl; 5,10-dihydroindeno[1,2-b]indol-10-yl; N-methyl- or N-phenyl-5,10-dihydroindeno [1,2-b]indol-10-yl; 5,6-dihydroindeno[2,1-b]indol-6-yl; N-methyl-or N-phenyl-5,6-dihydroindeno[2,1-b]indol-6-yl; azapentalene-4-yl; thiapentalene-4-yl; thiapentalene-6-yl; thiapentalene-6-yl; mono-, di- and tri-methyl-azapentalene-4-yl.
- 8. The catalyst system according to claim 1 wherein, in said metallocene complex of formula (I), L is selected from the group consisting of -Cl, -Br, -Me, -Et, -n-Bu, -sec-Bu, -Ph, -Bz, -CH₂SiMe₃, -OEt, -OPr, -OBu, -OBz and -NMe₂.

9. The catalyst system according to claim 1 wherein, in said organometallic aluminium compound of formula (II), Ar corresponds to formula (III):

$$R^{6}$$
 R^{7}
 R^{8}
 R^{9}
(III)

wherein R^6 , R^8 and R^{10} are selected from the group consisting of hydrogen, halogen, $-R^3$, $-C(O)R^3$, $-OR^3$, $-SR^3$, $-NR^3$ ₂ and $-NO_2$, wherein R^3 has the meaning reported in claim 1; the groups R^7 and R^9 are selected from the group consisting of hydrogen, halogen, linear or branched, saturated or unsaturated C_1 - C_{20} alkyl, C_3 - C_{20} cycloalkyl, C_6 - C_{20} aryl, C_7 - C_{20} alkylaryl and C_7 - C_{20} arylalkyl groups, optionally containing one or more Si or Ge atoms; two adjacent substituents R^6 - R^{10} optionally form a ring, having 3 to 8 carbon atoms;

R⁴ is a C₁-C₅ alkyl group; and

R⁵ is hydrogen or a C₁-C₅ alkyl group.

- 10. The catalyst system according to claim 9, wherein Ar is selected from the group consisting of 4-fluoro-phenyl, 4-chloro-phenyl, 4-methoxyphenyl, 4-nitrophenyl, 3-methylphenyl, 3-isopropylphenyl, 2,4-difluorophenyl, 2,4-dichlorophenyl, 2,6-difluorophenyl, 3,5-dichlorophenyl, 2,4,6-trifluorophenyl, 3,5-dichlorophenyl, 3,4,5-trifluorophenyl, 3,4,5-trichlorophenyl, pentafluorophenyl and pentachlorophenyl.
- 11. The catalyst system according to claim 1, wherein said organometallic aluminium compound of formula (II) is selected from the group consisting of tris(2-phenyl-propyl)aluminium, tris[2-(4-fluoro-phenyl)-propyl]aluminium, tris[2-(4-chloro-phenyl)-propyl]aluminium, tris[2-(4-chloro-phenyl)-propyl]aluminium, tris(2-phenyl-butyl)aluminium, tris(3-methyl-2-phenyl-butyl)aluminium, tris(2-phenyl-pentyl)aluminium, tris[2-(pentafluorophenyl)-propyl]aluminium, tris[2,2-diphenyl-ethyl]aluminium and tris[2-phenyl-2-methyl-propyl]aluminium.
- 12. A catalyst system for the polymerization of olefins comprising the product obtained by contacting the following components:

(A) a metallocene complex of formula (I):

$$(Cp)(ZR_{m}^{1})_{n}(A)_{r}ML_{p}$$
 (I)

wherein M, Cp, $(ZR_m^1)_n$, A, L, r and p have the meanings reported in claim 1; and

(B') the product of the reaction between water and an organometallic aluminium compound of formula (II):

Al
$$[CH_2-C(Ar)R^4R^5]_{\star}H_{\star}$$
 (II)

wherein Ar, R⁴, R⁵, x and y have the meanings reported in claim 1;

the molar ratio between said organometallic aluminium compound and said water being comprised between 1:1 and 100:1.

13. The catalyst system according to claim 1 or 12, wherein said metallocene complex is pre-alkylated with one or more organometallic aluminium compounds of formula (IV):

$$AlR^{11}_{3-z}H_z$$
 (IV)

wherein R^{11} is a linear or branched, saturated or unsaturated, C_1 - C_{10} alkyl, C_6 - C_{20} aryl, C_7 - C_{20} arylalkyl or C_7 - C_{20} alkylaryl and z is 0 or 1.

14. An alumoxane obtained by contacting an organometallic aluminium compound of formula (II)

Al
$$[CH_2-C(Ar)R^4R^5]_xH_v$$
 (II)

wherein Ar, R⁴, R⁵, x and y have the meanings reported in claim 1, with water, the molar ratio between said organometallic aluminium compound and said water being comprised between 1:1 and 100:1.

- 15. A process for the polymerization of one or more olefins in the presence of a catalyst system according to any of claims 1 to 13.
- 16. The process according to claim 15, wherein said olefin is an α -olefin of formula CH₂=CHR, wherein R is hydrogen or a C₁-C₂₀ alkyl radical.
- 17. The process according to claim 16, wherein said α -olefin is selected from the group consisting of propylene, 1-butene, 4-methyl-1-pentene, 1-hexene and 1-octene.
- 18. The process according to claim 15, wherein ethylene is copolymerized with an α-olefin of formula CH₂=CHR', wherein R' is a linear, branched or cyclic C₁-C₂₀ alkyl radical, or with a cycloolefin, and optionally with a polyene.

(19) World Intellectual Property Organization International Bureau



(43) International Publication Date 29 March 2001 (29.03.2001)

PCT

(10) International Publication Number WO 01/21674 A1

(51) International Patent Classification7: C08F 10/00, 4/64

(21) International Application Number: PCT/EP00/09111

(22) International Filing Date:

15 September 2000 (15.09.2000)

(25) Filing Language:

English

(26) Publication Language:

English

(30) Priority Data: 99203110.4 22 September 1999 (22.09.1999) EF

(71) Applicant (for all designated States except US): BASELL TECHNOLOGY COMPANY B.V. [NL/NL]; Hocksteen 66, NL-2132 MS Hoofddorp (NL).

(72) Inventors; and

(75) Inventors/Applicants (for US only): VAN BAAR, Jan, F. [NL/NL]; De Terp 45, NL-1911 AB Uitgeest (NL). HORTON, Andrew, D. [GB/NL]; Tituaanstraat 1, NL-1077 RC Amsterdam (NL). VAN KESSEL, Gerard, M., M. [NL/NL]; Koningsspil 6, NL-4191 TB Geldermalsen

(NL). KRAGTWIJK, Eric [NL/NL]; Hilversumstraat 30, NL-1024 JZ Amsterdam (NL). VAN DE PAS, Max [NL/NL]; Zwart Janstraat M5a, NL-3035 AP Rotterdam (NL). SCHUT, Peter, A. [NL/NL]; Forelstraat 26, NL-1317 PT Almere (NL). STAPERSMA, Johan [NL/NL]; Heereweg 7-b, NL-1901 MA Castricum (NL).

(74) Agents: COLUCCI, Guiseppe et al.; Basell Poliolefine Italia S.p.A., Intellectual Property, P. le G. Donegani, 12, I-44100 Ferrara (IT).

(81) Designated States (national): AU, BR, CA, CN, CZ, HU, ID, IL, IN, JP, KR, MX, NO, PL, RU, SG, US, ZA.

(84) Designated States (regional): European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE).

Published:

With international search report.

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

(54) Title: CATALYST SYSTEM AND PROCESS FOR THE POLYMERIZATION OF OLEFINS

(57) Abstract: A catalyst system for the polymerization of olefins comprising the product obtained by contacting. (A) a metal-locene complex, (B) an organometallic aluminium compound of formula (II): A1[CH₂-C(Ar)R⁴R⁵], H_y wherein Ar is a substituted or unsubstituted aryl group having from 6 to 20 carbon atoms; R⁴ is a linear or branched, satured or unsaturated, C_1 - C_{10} alkyl. C_6 - C_{20} aryl, C_7 - C_{20} arylalkyl or C_7 - C_{20} alkylaryl; R⁵ is hydrogen or a linear or branched, saturated or unsaturated, C_1 - C_{10} alkyl. C_6 - C_{20} aryl, C_7 - C_{20} arylalkyl or C_7 - C_{13} alkylaryl group; x = 2 or 3; y = 3-x: and (C) water; the molar ratio (B)/(C) being comprised between 1:1 and 100:1. These catalysts show an improved activity with respect to known catalysts, wherein different aluminium compounds are used.

se type a plus sign (+) inside	this box>	Appr	oved for use the	ough 9/30/00 O	SB/01 (12-97) MB 0651-0032	4			
Under the Paperwork Red a valid OMB control numb	luction Act of 1995, no person	Patent and Trademarks are required to respond	k Office; U.S. E to a collection	DEPARTMENT OF Information unl	F COMMERCE ess it contains	•			
		Attorney Docket	t Number	US 19005	5	\geq			
	FOR UTILITY OR SIGN	First Named Inv	entor	Jan F. V	/an Baar	e			
	PLICATION	co	MPLETE IF	KNOWN					
	R 1.63)	Application Num	Application Number						
(27 23	,	Filing Date	М	arch 19,	2002				
Declaration [Submitted OR	☐ Declaration Submitted after Initia	al Group Art Unit							
with Initial Filing	Filing (surcharge (37 CFR 1.16 (e)) required)	Examiner Name				\neg			
	requireu)								
the specification of which is attached hereto OR was filed on (MM/D Application Number PCT). I hereby state that I have reamended by any amendment I acknowledge the duty to continue the propertificate, or 365(a) of any America, listed below and hear of any PCT international according to the specific state of the specific state o	DD/YYYY) 09/15/200/EP00/0911 and wa eviewed and understand the cent specifically referred to abordisclose information which is not the property of the propert	as Unite as amended on (MM/DD/Y contents of the above identive anaterial to patentability as 119(a)-(d) or 365(b) of an which designated at least before that of the application	d States Applic YYY) tified specificate defined in 37 C my foreign application ign application ign application on which p	ation Number or F on, including the c FR 1 56 ication(s) for pate other than the t for patent or inversionally is claimed	(if applicable) claims, as ent or inventor's United States of ntor's certificate,	, 			
Prior Foreign Application Number(s)	Country	Foreign Filing Date (MM/DD/YYYY)	Priority Not Claimed		opy Attached?				
99203110.4	EP	09/22/1999	000						
Additional foreign applic	ation numbers are listed on a	supplemental priority data	sheet PTO/SB	/02B attached he	reto.	_			
I hereby claim the benefit in Application Number		(MM/DD/YYYY)	i application(s)	listed below		\dashv			
Application Number	rung Date	. (1111)	num	gn application (s) for patent or inventor's country other than the United States of lication for patent or inventor's certificate, which priority is claimed Certified Copy Attached?					

-- .*- --

[Page 1 of 4]

Burden Hour Statement This form is estimated to take 0.4 hours to complete Time will vary depending upon the needs of the individual case Any comments on the amount of time you are required to complete this form should be sent to the Chief Information Officer, Patent and Trademark Office, Washington, DC 20231 DO NOT SEND FEES OR COMPLETED FORMS TO THIS ADDRESS. SEND TO: Assistant Commissioner for Patents, Washington, DC 20231

Please	Under	+) inside this box → the Paperwork Redu OMB control numbe	ection Act of 1995, no	F o persons are	atent and Tra required to r	demark Offi	ce. US [DEPART	30/00 OMB MENT OF CO	DMMERC	2 T	_
	DECLA	RATION	— Utility	or D	esign	Pate	nt A	\ppl	icatio	n		
United United Informa	States of Americ States or PCT Int tion which is mai	it under 35 U S C 13 a, listed below and, ernational application terial to patentability international filing da	insofar as the subj n in the manner prov as defined in 37 Cl	ect matter of vided by the f FR 1 56 which	each of the irst paragrapl	claims of th of 35 U S (is applica C 112, La	ition is no acknowle	ot disclosed dge the duty	in the pric to disclos	or se	
	U.S. Pare	nt Application Number	or PCT Parent		Parent Filing Date (MM/DD/YYYY)			Parent Patent Number (if applicable)				
PC	T/EP00/0	9111			09/15/20	000						
☐ Ac	lditional U.S. or F	CT international app	olication numbers are	e listed on a	upplemental	priority data	sheet PT	O/SB/02I	B attached he	ereto	1	
			Customer Numi	oer				•	all business i Place Custo Number Bar Label her	mer Code	∌nt	
	Nam		Regist	ration	name/registration number listed below Name				Registration Number			
/	anne W. F Lliam R.	Patterson Reid	31,21 47,89									
	all corresponde		stomer Number Bar Code Label	Registered F	ractitioner Inf	7	_		dence addr		w	
Addr	Rac	sell USA I	nc.							-	_	
Addr	013	Appleton	Road					•				
City	Ell	ton_			State	MD	ZIP	2	1921		_	
Coun			Telephor		996-165		1 (4.7)		996-15		_	
believe punish:	d to be true; and	d further that these nprisonment, or bot	are true and that all statements made on information and belief are the knowledge that willful false statements and the like so made are d that such willful false statements may jeopardize the validity of the						re			
) Name	Name of Sole or First Inventor:					A petition has been filed for this unsigned inventor						
		Given Name (first and middle [if any])				Family Name or S				Surname		
Inver	JAN-E.,	<u> </u>	80		VAN BAAR		<u> </u>				\dashv	
Signa	ature	Amatardam		aal	TAIR				Date	Feb NL	<u>2</u> 6,	2
Resid	ence: City	Amsterdam	State		Country	NL			Citizenship	1 76-	\dashv	
Post	Office Address	De Terp, 45									\dashv	
Post	Office Address	A	1	<u> </u>	1911 A		,	—т	NL		\dashv	
City		Amsterdam so	tate	ZiP			Cou		B/02A attac		\dashv	

[Page 2 of **4**]

Control of the Book of the Boo

. Р) inside this box → r the Paperwork Reduction Act OMB control number	of 1995, n	o persons a	Patent and are required to	Trademark Office	e; US DE	igh 9/30/98 PARTMENT	O/SB/02A (3-9 OMB 0651-00 OF COMMERO Iless it contains	032 _ CE		
	D	DECLARATION				ADDITIONAL INVENTOR(S) Supplemental Sheet Page of						
5-00	Name of Addition	nal Joint Inventor, if an	ıy:		A petiti	on has been file	ed for this	unsigned i	nventor	7		
	Given Na	me (first and middle [if any]])	Family Name or S					Surname			
	MAX	_	1		VAN D	E PAS						
	Inventor's Signature	His			, and service			Date	Feb 26	, 2		
	Residence: City	Rotterdam	State	NLX	Country	NL		Citizenship	NL	_		
	Post Office Address	Zwart Janstraat, 11	5 a									
	Post Office Address		1 1							_		
	City	Rotterdam	State		ZIP	3035 AP	Country	NL	·	_		
<i>a</i> : A	Name of Addition		A petition has been filed for this unsigned inventor									
-00	Given Name (first and middle [if any])				Family Name or Surname							
-	PETER A.	1-			sc	HUT				1		
-	Inventor's Signature	Affr				,		Date	Feb 2	6,		
	Residence: City	Almere	State	NL	Country	NL		Citizenship	NL			
	Post Office Address	Forelstraat, 26]		
	Post Office Address			·								
	City	Almere	State		ZIP	1317 PT	Countr	y NL				
• •	Name of Addition	nal Joint Inventor, if an	y:		A petition	on has been file	d for this	unsigned i	nventor			
	Given Name (first and middle [if any])				Family Name or Surname							
ω	JOHAN				STAPERSMA							
	Inventor's Signature		81					Date	Feb 26	, 2		
	Residence: City	Castricum	State	NN	Country	NL		Citizenship	NL			
	Post Office Address	Heereweg, 7-b]		
	Post Office Address									_		
	City	Castricum	State		ZIP	1901 MA	Cou	untry NL				

		ECLARATIO	N		ADDITIONAL INVENTOR(S) Supplemental Sheet Page 3_ of 4_					
2-00	Name of Addition	Name of Additional Joint Inventor, if any:				on has been filed f	or this	unsigned i	nventor	_
	Given Na	ne (first and middle [if any])				Family Name	or Su	ırname		
	ANDREW	D.								
	Inventor's Signature	A.D.	Hor	W.				Date	Feb	26,
	Residence: City	Amsterdam	State	NLX	Country	NL		Citizenship	GB	
	Post Office Address	Titiaanstraat, 1								
	Post Office Address									
	City	Amsterdam	State		ZIP	1077 RC _{co}	untry	Amster	dam	
20	Name of Additional Joint Inventor, if any:									
00	Given Name (first and middle [if any])				Family Name or Surname					
	GERARD M.M.				VAN KESSEL					
	Inventor's Signature	grae						Date	Feb	26,
	Residence: City	Geldermalsen	State	NLX	Country	NL		Citizenship	, NL	
	Post Office Address	Koningsspil, 6								
	Post Office Address									
	City	Geldermalsen	State		ZIP	4191 TB	Count	_{ry} NL		
00	Name of Additio	nal Joint Inventor, if a	ny:		A petition	on has been filed f	or this	unsigned i	nventor	
	Given Name (first and middle [if any])				Family Name or Surname					
	ERIC		K	KRAGTWIJK						
	Inventor's Signature	Ene 1	Ria	this	h			Date	Fel	26
	Residence: City	Amsterdam_	State	NLX	Country	NL		Citizenship	, NL	
	Post Office Address	Hilversumstraat, 30								
	Post Office Address			_		_				
	City	Amsterdam			ZIP	1024 JZ		untry NL		